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(54) Tide: I.PA RECEPTOR AGONISTS AND ANTAGONISTS AND METHODS OF USE

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WO 01/71022 A2 (57) Abstract: The present invention relates to compounds according to formula (f) as disclosed herein as well as pharmaceutical compositions which broulds those compounds. Also disclosed are methods of using such compounds, which have sativity as agondsts or as autgentists of LPA receptor, such methods including inhibiting LPA entirely on an LPA receptor, modulating LPA receptor, reading cancer, onhancing cell proliferation, and treating a wound.

> WO 01/71022 PCT/US01/08729

LPA RECEPTOR AGONISTS AND ANTAGONISTS AND METHODS OF USE

reference in its entirety. Serial No. 60/190,370 filed March 17, 2000, which is hereby incorporated by This application claims benefit of U.S. Provisional Patent Application

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FIELD OF THE INVENTION

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therapeutic uses thereof including, but not limited to, prostate cancer therapy, ovarian cancer therapy, and wound healing. which bave activity as either agonists or antagonists on LPA receptors and various This invention relates to lysophosphatidic acid ("LPA") derivatives

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BACKGROUND OF THE INVENTION

phosphate, sphingosylphosphorylcholine (SPC), and sphingosine (SPH). Exemplary SPMs include sphingosine-1-phosphate (SPP), dihydrosphingosine-1contains the sphingolipid mediators (SPMs), which possess a sphingoid base motif phospholipid growth factors (PLGFs). In spite of their similar pharmacologic phosphate (alkenyl-GP), and lysophosphatidyl serine (LPS). The second category phosphatidic acid (PA), cyclic phosphatidic acid (cyclic-PA), alkenyl glycerol (GPMs), which possess a glycerol backbone. Exemplary GPMs include LPA, two broad categories. The first category contains the glycerophospholipid mediators Tokumura, 1995; Moolenzar et al., 1997). PLGFs can be sub-divided structurally into properties in inducing the proliferation of most quiescent cells (Jalink et al., 1994a; with growth factor-like properties has been discovered, collectively known as proliferation. In addition to polypeptide growth factors, an emerging class of lipids All non-transformed cells require growth factors for their survival and

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PLGFs present in the serum and plasma that exhibit growth factor-like properties: lipid mediators have been identified and characterized. There are still, yet unknown, (Yatomi et al., 1995), and SPC (Tigyi et al., 2000) have been detected in serum. These 1989), alkenyl-GP (Liliom et al., 1998), cyclic-PA (Kobayashi et al., 1999), SPP LPA (Tigyi et al., 1991; Tigyi and Miledi, 1992), PA (Myher et al.

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Tigyi and Miledi, 1992). LPA, with its <20 µM concentration, is the most abundam PLGF present in the serum (Tigyi and Miledi, 1992; Jalink et al., 1993). In eukaryotic cells, LPA is a key intermediate in the early stages of phospholipid biosynthesis, which takes place predominantly in the membrane of endoplasmic reticulum (ER) (Bosch, 1974; Bishop and Bell, 1988). In the ER, LPA is derived from the action of Acyl-CoA on glycerol-3-phosphate, which is further acylated to yield PA. Because the rate of acylation of LPA to PA is very high, very little LPA accumulates at the site of biosynthesis (Bosch, 1974). Since LPA is restricted to the ER, its role as a metabolic intermediate is most probably unrelated to its role as a signaling molecule.

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LPA is a constituent of serum and its levels are in the low micromolar (µM) range (Eicholtz et al., 1993). This level is expected because LPA is released by activated platelets during the coagulation process. Unlike serum, it is not detectable in fresh blood or plasma (Tigyi and Miledi, 1992; Eicholtz et al., 1993). LPA that is present in the serum is bound to abumin, and is responsible for a majority of the heat stable, and non-dialysable biological activity of the whole serum (Moolenaar, 1994). The active serum component that is responsible for eliciting an inward chloride current in Xenopus oocyte was indentified to be LPA (18:0) (Tigyi and Miledi, 1992). The bulk of the abumin-bound LPA(18:0) is produced during the coagulation process, rather than by the action of lysophospholipase D (PLD) on lyso-PC. The latter pathway is responsible for the presence of LPA in 'aged' plasma that has been decoagulated by the action of heparin or citrate plus dextrose (Tokumura et al., 1986). Another point to note is that LPA is not present in plasma that has been treated with EDTA. This fact implies that plasma lysophospholipase may be Ca²*-dependent (Tokumura et al., 1986).

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The role of albumin is to protect LPA from the actions of phospholipases present in the serum (Tigyi and Miledi, 1992). Tigyi and Miledi suggested that albumin not only acts as a carrier of LPA in the blood stream, but also increases its physiological half-life. There are yet unidentified lipid mediators present in serum albumin that mimic the actions of LPA in eliciting chloride current in *Konopus* oocyte.

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LPA-responsive cell types extend from slime mold amoebae and Xenopus oocyte to mammalian somatic cells. Thus, it seems likely that the source of LPA and its release may not be restricted only to activated platelets. Recent experiments showed that, on stimulation by peptide growth factors, mammalian fibroblasts rapidly produce LPA, which is followed by its release into the extracellular medium (Fukami and Takenawa, 1992).

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WO 01/71022

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PCT/US01/08729

There is evidence that relatively high amounts of bioactive LPA of unknown cellular origin are present in the ascitic fluid of ovarian cancer patients (Xu et al., 1995a), and that the ascitic fluid from such patients is known to possess potent mitogenic activity for ovarian carcinoma cells (Mills et al., 1988; Mills et al., 1990). It remains to be established whether it is secreted by tumor cells into the extracellular fluid, secreted by leukocytes, or produced from more complex lipids via the actions of various phospholipases.

GPMs and SPMs elicit a wide variety of cellular responses that span the phylogenetic tree (falink et al., 1993a). LPA induces transient Ca²⁺ signals that originate from intracellular stores in a variety of cells such as neuronal (falink et al., 1993; Durieux et al., 1992), platelets, normal as well as transformed fibroblasts (falink et al., 1999), epithelial cells (van Corven et al., 1989; Moolenaar, 1991), and *Kenopus* oocytes (Tigyi and Miledi, 1992; Durieux et al., 1989; Moolenaar, 1991), and *Kenopus* oocytes (Tigyi and Miledi, 1992; Durieux et al., 1979; Tokumura et al., 1979; Simon et al., 1982) and smooth muscle contraction (Tokumura et al., 1980; Tokumura et al., 1994), and upon intravenous administration it induces species-dependent changes in blood pressure ((Schumaeher et al., 1979; Tokumura et al., 1978).

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effects of LPA do not require the presence of peptide growth factors. This observation neurite retraction, which was accompanied by rapid, but transient, rounding of the cell and cell division (van Corven et al., 1989; van Corven et al., 1992). The growth-like methyl xanthin, the antimitogenic actions of LPA in Sp² myeloma cells were additive insulin or epidermal growth factor (Moolenaar, 1991) to sustain cell proliferation. A point to note is that, in Sp2 myleoma cells, LPA was responsible for an antimitogenic include formation of focal adhesions and stress fibers in fibroblasts (Ridley and Hall, LPA, when added to quiescent fibroblasts, stimulates DNA synthesis makes LPA different from endothelin or vasopressin, which require the presence of Fischer et al., 1998). Unlike the mitogenic pathway, the antimitogenic pathway was differentiation by inducing the retraction of developing neurites (Jalink et al., 1994a ; Moolenaar, 1992) to serum-starved NIE-115 neuroblastoma cells caused immediate not affected by pertussis toxin (PTX). Also, on addition of forskolin and isobutyl (Tigyi et al., 1994). In various cell types, LPA causes cytoskeletal changes, which neuroblastoma cells maintain their undifferentiated phenotype, but fail to undergo response, which was mediated by an increase in cAMP levels (Tigyi et al., 1994; Jalink et al., 1994b). Addition of nanomole (nmol) amounts of LPA (Jalink and body (Jalink et al., 1993b). When a continuous presence of LPA is provided, 1992). LPA also promotes the reversal and suppression of neuroblastoms

WO 01/71022 -4-PCT/US01/08729

morphological differentiation, the addition of LPA reverses this morphological mitosis (Jalink et al., 1993b). Additional factors, such as insulin-like growth factors, actin-cytoskeleton, rather than from loss of adhesion to the substratum (Jalink et al. change. Thus, LPA-induced neurite retractions result from the contraction of the were required for the progression of the cell cycle. Once the cells have undergone 1993b; Jalink et al., 1994b).

block neonatal cardiomyocyte apoptosis (Umansky et al., 1997). enhanced cell motility and increased cell adhesion. Finally, LPA is also known to hepatoma and carcinoma cells into the monolayer of mesothelial cells (Imamura et al. al., 1995). In addition to inducing cell migration, LPA promotes the invasion of induces cell migration by a haptotactic mechanism in human monocytes (Zhou et 1993). The mechanism that underlies this invasion is still unclear, but it may be due to LPA, similar to other physiological chemoattractants (e.g., interleukin

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responsible for cellular actions that were similar to or opposite to other GPMs, antiproliferative actions, unlike LPA, which induces proliferation. al., 1998). Murakami-Murofushi et al. (1993) showed that cyclic-PA exhibited current just like other GPMs; but its response was not desensitized by LPA (Fischer et depending on the cell type. When tested on the Xenopus oocyte, it elicited chloride A unique natural phospholipid, namely cyclic-PA, was shown to be

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and others is shown in Figure 1. their cellular responses via interacting with the heterotrimeric G-protein. A number of guanine nucleotide-binding regulatory protein (O protein)-coupled receptors (GPCR) and PSP-24. A phylogenetic tree illustrating the relatedness of these LPA receptors LPA receptors have been identified including, among others, EDG-2, EDG-4, EDG-7, superfamily. Seven-TM GPCRs are a family of cell-surface receptors that mediate PLGF receptors (PLGFRs) belong to a seven-transmembrane (7 TM)

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weight of 41 kDa (364 amino acids). Vzg-1 was very similar to an unpublished sheep expressed in cortical neurogenic regions and encoded a protein with a molecular highest expression in the brain and heart. rec1.3 (Macrae et al., 1996). It was widely distributed in the mouse tissue, with the also isolated as an orphan receptor from mouse and bovine libraries, and was known as sequence termed endothelial differentiation gene-2 (EDG-2). The same cDNA was encoding a putative serpentine receptor from mouse neocortical cell lines (Hecht et al., 1996). The gene was termed as ventricular zone gene-1 (Vzg-1). The gene was In 1996, Hecht et al. used differential hybridization to clone a cDNA

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putative LPA receptor PSP-24 (372 amino acids) from Xenopus oocyte (Guo et al., In 1996, Guo et al., using a PCR base protocol, isolated another 35

WO 01/71022 5 PCT/US01/08729

5 in heart tissue. of the EDG-2 human LPA receptor, led to two closely related GPCRs, namely, rat Northern analysis showed a high expression of mRNA that encoded EDG-3 and EGD 1996). A sequence based search for sphingolipid receptors, using the cDNA sequence H218 (EDG-5, 354 amino acids) and EDG-3 (378 amino acids) (An et al., 1997a). 1996). This receptor showed little similarity with Vzg-1/EDG-2/rec1.3 (Guo et al.,

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designated EDG-4 (An et al., 1998a). Northern blot analysis showed that, although leukocytes and testes (An et al., 1998a). different. Unlike EDG-2, EDG-4 was primarily expressed in peripheral blood prompted An et al. to perform a sequence-based search for a novel subtype of LPA EDG-2 and EDG-4 both serve as GPM receptors, their tissue distributions were very receptor (An et al., 1998a). A human cDNA, encoding a GPCR, was discovered and The recent identification of EDG-2 as a functional receptor for LPA

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previously unknown GPCR that belongs to the EDG family. The identified GPCR was designated EDG-7. It has a molecular mass of 40 kDa (353 amino acids). individual PLGFRs (Figure 1). The list continues to grow. two distinct families of PLOFs receptors PSP24 and EDG; with a total of ten expressed in heart, pancreas, prostate, and testes (Bandoh et al., 1999). Thus, there are Northern blot analysis of EDG-7 expression in human tissues showed that it is PCR amplification cDNA from human Jurkat T cells identified a

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specificities for GPMs or SPMs, as shown in Table 1 below These various receptors can be classified based on their ligand 20

Table 1: Phospholipid Growth Factor Receptor, Length and Principle Ligand
PLGFR Number of amino acids Principle Ligan EDG-3 EDG-2 EDG-8 EDG-7 EDG-6 EDG-4 EDG-1 385 353 400 372 373 354 382 378 364 SPP LPA LPA LPA L₽Α LΡΑ ¥3 SPP SPP SPP

Fischer et al., 1998) evoked oscillatory chloride-currents. These are not structurally Xenopus PSP24 and murine expressed PSP24 specifically transduce GPM (LPA, 25

Kenopus PSP24 Murine PSP24

- 6 -

homologous to the EDG family (Tigyi and Miledi, 1992; Fernhout et al., 1992). The EDG family can be divided into two distinct subgroups. The first group includes EDG-2, EDG-4, and EDG-7, which serve as receptors for only GPM (Hecht et al., 1996; An et al., 1998b) and transmit numerous signals in response to ligand binding. The second group involves EDG-1, EDG-3, EDG-5, EDG-6, and EDG-8, and is specific for SPMs (An et al., 1997a; Im et al., 2000; van Brocklyn et al., 1998; van Brocklyn et al., 2000; Spiegel and Milstein, 2000). Principle tissue expression of the various PLGFR's is shown in Table 2 below.

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Table 2: Human Tissue Expression of Phospholipid Growth Factor Receptors

Human Tissue with Highest Expression	Ubiquitous	Cardiovascular, CNS, Gonadal tissue, GI	Cardiovascular, Leukocyte	Leukocyte, Testes	Cardiovascular, CNS, Gonadal tissue, Placenta	Lymphoid, Hematopoietic tissue	Heart, Pancreas, Prostate, Testes	Brain	970
PLGFR	EDGI	EDG-2	EDG-3	EDG-4	EDG-5	EDG-6	FDG-7	BDG-8	PSP24

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PLGFs activate multiple G-protein-mediated signal transduction events. These processes are mediated through the heterotrimeric G-protein families G_{0/11}, G₁₀, and G₁₂₁₃ (Moolenaar, 1997, Spiegel and Milstein, 1995; Gohla, et al., 1998).

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The G₉11 pathway is responsible for phospholipase C (PLC) activation, which in turn induces inositol triphosphate (B₃) production with subsequent mobilization of Ca^{2*} in a wide variety of cells (Tokumura, 1995). In some cells, this response is PTX-sensitive, implying that there is involvement of multiple PTX-sensitive and insensitive pathways (Tigi et al., 1996). This pathway is also responsible for the diacyl glycerol (DAG)-mediated activation of protein kinase C (PKC). PKC activates cellular phospholipase D (PLD), which is responsible for the hydrolysis of phosphatidyl choline into free choline and PA (van der Bend et al., 1992a). Also, PLC is capable of activating MAP kinase directly, or via DAG activation of PKC in some cell types (Ghosh et al., 1997).

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The mitogenic-signaling pathway is mediated through the G-protein heterotriment G_{P0} subunit. Transfection studies indicate that the G_{P0} dimer rather than the α subunit is responsible for Ras-MAP kinase activation. The activation of

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WO 01/71022

-7-

PCT/US01/08729

phosphorylates and descinsitizes the receptor. The phosphorylated receptor is recruited 1999). The transactivated RTKs, in turn, activate Ras, which leads to the activation of ransactivated RTKS activate Ras, which leads to the activation of MAP kinases (ERK (2) via Raf. The Gia subunit, which is PTX-sensitive, inhibits adenylyl cyclase (AC), Ras is preceded by the transactivation of the receptor tyrosine kinases (RTKs) such as 11., 1989; van Corven et al., 1992), whereas antimitogenesis is accompanied by a nonthrough the Gia pathway, which results in decreased levels of cAMP (van Corven et MAP kinases (ERK 1,2) via Raf. The Gra subunit, which is PTX-sensitive, inhibits PTX sensitive Ca2+-dependent elevation of cAMP (Tigyi et al., 1994; Fischer et al., generating its active conformation (Lin et al., 1997; Ahn et al., 1999; Luttrell et al., opposing effects on the cAMP second messenger system. Mitogenesis is mediated resulting in By dimer docking to a G-protein-coupled receptor kinase (GRKs) that by B-arrestin, thus recruiting src kinase, which phosphorylates the EGF-receptor, AC, resulting in decreased levels of cyclic-AMP (cAMP). The opposite cellular effects by LPA, that is, mitogenesis and antimitogenesis, are accompanied by EGF (Cunnick et al., 1998) or PDGF receptors (Harrlich et al., 1998). The

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ibers (Ridley and Hall, 1992; Gonda et al., 1999), stimulation of chemotaxis (Jalink et that lead to cellular effects like retraction of neurites (Tigyi and Miledi, 1992; Tigyi et .999; Luttrell et al., 1999; Gohla et al., 1998) and converge on a small GTPase, Rho pecifically ribosylates Rho in an ADP-dependent mechanism (Imamura et al., 1996). ul., 1996; Dyer et al., 1992; Postma et al., 1996; Sato et al., 1997), induction of stress signaling pathway, which leads to the rearrangement of the actin-cytoskeleton. This ccumulation of the phosphorylated form of MLC, leading to cytoskeletal responses avasiveness (Imamura et al., 1993; Imamura et al., 1996). The PLGF-induced, Rhopathway may also involve the transactivation of RTKs (Lin et al., 1997; Ahn et al., Moolenaar, 1997). Much more is known about the down-stream signaling of Rho al., 1993a), cell migration (Zhou et al., 1995; Kimura et al., 1992), and tumor cell occause various protein partners have been isolated and identified. Rho activates Ser/Thr kinases, which phosphorylate, and as a result inhibit, myosin light chain phosphatase (MLC-phosphatase) (Kimura et al., 1996). This path results in the In contrast, very little is known about the PTX-insensitive G1211 nediated, tumor cell invasiveness is blocked by C. Botulinium C3-toxin, which

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Rho also has the ability to stimulate DNA synthesis in quiescent fibroblasts (Machesky and Hall, 1996; Ridley, 1996). The expression of Rho family GTPase activates serum-response factor (SRF), which mediates early gene transcription (Hill et al., 1995). Furthermore, PLGF (LPA) induces tumor cell

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WO 01/71022 -8-

invasion (Imamum et al., 1996); however, it is still unclear whether it involves cytoskeletal changes or gene transcription, or both.

By virtue of LPA/LPA receptor involvement in a number of cellular pathways and cell activities such as proliferation and/or migration, as well as their implication in wound healing and cancer, it would be desirable to identify novel compounds which are capable of acting, preferably selectively, as either antagonists or agonists at the LPA receptors identified above.

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There are currently very few synthetic or endogenous LPA receptor inhibitors which are known. Of the antagonists reported to date, the most work was done on SPH, SPP, N-palmitoyl-1-serine (Bitman et al., 1996), and N-palmitoyl-1-tyrosine (Bitman et al., 1996). It is known that the above-mentioned compounds inhibit LPA-induced chloride currents in the Xenopus oocyte (Bitman et al., 1996; Zsiros et al., 1998). However, these compounds have not been studied in all cell systems. It is also known that SPP inhibits tumor cell invasiveness, but it is uncertain whether SPP does so by being an inhibitor of LPA or via the actions of its own receptors. N-palmitoyl-1-serine and N-palmitoyl-1-tyrosine also inhibited LPA-induced platelet aggregation (Sugiura et al., 1994), but it remains to be seen whether these compounds act at the LPA receptor. Lysophosphatidyl glycerol (LPG) was the first lipid to show some degree of inhibition of LPA actions (van der Bend et al., 1995). None of these inhibitors was shown to selectively act at specific LPA receptors.

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A polysulfonated compound, Surarnin, was shown to inhibit LPA-induced DNA synthesis in a reversible and dose-dependent manner. However, it was shown that Suramin does not have any specificity towards the LPA receptor and blocked the actions of LPA only at very high millimolar (mM) concentrations (van Corven et al., 1992).

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The present invention is directed to overcoming the deficiencies associated with current LPA agonists and LPA antagonists.

SUMMARY OF THE INVENTION

The present invention relates to compounds according to formula (I) as

follows:

WO 01/71022

PCT/US01/08729

-9-

PCT/US01/08729

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wherein

at least one of X¹, X², and X³ is (HO)₂PO—Z¹— or (HO)₂PO—Z²—P(OH)O—Z¹—, X¹ and X² are linked together as —O—PO(OH)—O—, or X¹ and X² are linked together as —O—PO(OH)—NH—;

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at least one of X^1 , X^2 , and X^3 is R^1-Y^1-A- with each being the same or different when two of X^1 , X^2 , and X^3 are R^1-Y^1-A- , or X^2 and X^3 are Rinked together as $-N(H)-C(O)-N(R^1)-$;

optionally, one of X^1 , X^2 , and X^3 is H;

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A is either a direct link, $(CH_2)_k$ with k being an integer from 0 to 30, or O;

 Y^1 is $-(CH_2)$ — with I being an integer from 1 to 30, -O—,

15 —C—,—S—, or—NR²—;

 Z^1 is $-(CH_2)_{m}$ — or $-O(CH_2)_{m}$ — with m being an integer from 1 to 50, $-C(R^3)H$ —, -NH—, -O—, or -S—;

 Z^2 is $-(CH_2)_n$ or $-O(CH_2)_n$ with n being an integer from

1 to 50 or —O—;

 Q^1 and Q^2 are independently $H_{2s} = NR^4$, = 0, or a combination of H and $-NR^2R^6$,

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R¹, for each of X¹, X², or X³, is independently hydrogen, a straight or branched-chain C1 to C30 alkyl, a straight or branched-chain C2 to C30 alkenyl, an aromatic or heteroaromatic ring with or without mono-, di-, or trisubstitutions of the ring, an acyl including a C1 to C30 alkyl or an aromatic or heteroaromatic ring, an arylatkyl including straight or branched-chain C1 to C30 alkyl, an aryloxyalkyl including straight or branched-chain C1 to C30 alkyl,

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PCT/US01/08729

beteroaromatic ring, an arylalkyl including straight or branched-chain C1 to C30 alkyl, R2, R3, R4, R5, R6, R7, and R8 are independently hydrogen, a straight or branched-chain C1 to C30 alkyl, a straight or branched-chain C2 to C30 alkenyl, an aromatic or heteroaromatic ring with or without mono-, di-, or trisubstitutions of the ring, an acyl including a C1 to C30 alkyl or aromatic or or an aryloxyalkyl including straight or branched-chain C1 to C30 alkyl;

acid, phosphatidic acid, cyclic phosphatidic acid, alkenyl glycerolphosphate, dioctyl wherein the compound of formula I is not lysophosphatidic glycerol pyrophosphate, or N-palmitoyl-L-serine.

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Also disclosed are pharmaceutical compositions which include a pharmaccutically-acceptable carrier and a compound of the present invention.

an LPA receptor with the compound under conditions effective to inhibit LPA-induced the present invention which has activity as an LPA receptor antagonist and contacting inhibiting LPA activity on an LPA receptor which includes providing a compound of A further aspect of the present invention relates to a method of activity of the LPA receptor.

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modulating LPA receptor activity which includes providing a compound of the present invention which has activity as either an LPA receptor agonist or an LPA receptor antagonist and contacting an LPA receptor with the compound under conditions Another aspect of the present invention relates to a method of effective to modulate the activity of the LPA receptor

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administering an effective amount of the compound to a patient in a manner effective treating cancer which includes providing a compound of the present invention and Still another aspect of the present invention relates to a method of

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invention which has activity as an agonist of an LPA receptor and contacting the LPA receptor on a cell with the compound in a manner effective to enhance LPA receptor-Yet another aspect of the present invention relates to a method of enhancing cell proliferation which includes providing a compound the present induced proliferation of the cell,

A further aspect of the present invention relates to a method of treating compound to a wound site, where the compound binds to LPA receptors on cells that promote healing of the wound, thereby stimulating LPA receptor agonist-induced cell activity as an agonist of an LPA receptor and delivering an effective amount of the a wound which includes providing a compound of the present invention which has proliferation to promote wound healing.

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A still further aspect of the present invention relates to a method of making the compounds of the present invention. One approach for making the compounds of the present invention includes:

reacting
$$(Y^2O)_2PO-Z^{11}-Z^{13}$$
 or $(Y^2O)_2PO-Z^{12}-P(OH)O-Z^{11}-Z^{13}$,

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first leaving group; and

with an intermediate compound according to formula (VI) Y2 is H or a protecting group,

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at least one of X11, X12, and X13 is R11-Y11-A- with each being the same or different when two of X11, X12, and X13 are R11. Y11-A-, or X12 and X13 are linked together as -N(H)-C(O)-

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leaving group; A is either a direct link, $(CH_2)_k$ with k being an integer from 0 optionally, one of X11, X12, and X13 is H;

at least one of X11, X12, and X13 is OH, NH2, SH, or a second

Y¹¹ is —(CH₂)— with *l* being an integer from 1 to 30, —O—,

Hand -NR'R'S; Q¹ and Q² are independently H₂, =NR¹³, =O, a combination of

C2 to C30 alkenyl, an aromatic or heteroaromatic ring with or without straight or branched-chain C1 to C30 alkyl, a straight or branched-chain R¹¹, for each of X¹¹, X¹², or X¹³, is independently hydrogen, a

mono-, di-, or tri-substitutions of the ring, an acyl including a C1 to straight or branched-chain C1 to C30 alkyl, straight or branched-chain C1 to C30 alkyl, an aryloxyalkyl including C30 alkyl or an aromatic or heteroaromatic ring, an arylalkyl including

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mono-, di-, or tri-substitutions of the ring, an acyl including a C1 to including straight or branched-chain C1 to C30 alkyl; straight or branched-chain C1 to C30 alkyl, or an aryloxyalkyl C30 alkyl or aromatic or heteroaromatic ring, an arylalkyl including C2 to C30 alkenyl, an aromatic or beteroaromatic ring with or without straight or branched-chain C1 to C30 alkyl, a straight or branched-chain R¹², R¹³, R¹⁴, R¹⁵, R¹⁶, and R¹⁷ are independently hydrogen, a

(HO)2PO-Z2-P(OH)O-Z1according to formula (I) where one or two of X^1 , X^2 , and X^3 is $(HO)_h PO - Z^1$ or deprotection step being performed under conditions effective to afford a compound followed by a de-protection step, if necessary, with both said reacting and the չչ

WO 01/71022

-13-

PCT/US01/08729

to inhibit or enhance, respectively, biochemical pathways mediated by LPA receptor specific and substantial uses in treating cancer and enhancing wound healing. herein as being either agonists or antagonists of one or more LPA receptors find uses signaling. By modulating LPA receptor signaling, the antagonists and agonists find The compounds of the present invention which have been identified

BRIEF DESCRIPTION OF THE DRAWINGS

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EDG-2, EDG-4, EDG-7, and PSP-24 (α,β) relatedness of ten phospholipid growth factor receptors, including LPA receptors Figure 1 is a phylogenetic tree illustrating the classification and

serine amide compounds 35-43. Figure 2 illustrates the synthesis scheme employed for preparation of

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scrine amide phosphate compounds 55-59. Figure 3 illustrates the synthesis scheme employed for preparation of

biphosphate compounds 66-68. Figure 4 illustrates the synthesis scheme employed for preparation of

5A illustrates the synthesis scheme employed for preparation of 1,2-biphosphate biphosphate compounds. compounds 85-92. Figure 5B illustrates a synthesis scheme for preparing 1,3-Pigures 5A-B illustrate synthesis of biphosphate compounds. Figure

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pyrophosphate compounds. Figures 6A-B illustrate synthesis schemes for preparation of

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mono-phosphates and mono-phosphonates from a tosylate-protected di-ether Figures 7A-C illustrate synthesis schemes for preparation of substituted

straight-chain fatty acid phosphate compounds 106-110. Figure 8 illustrates the synthesis scheme employed for preparation of

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monoalkyi esters. Figure 9 illustrates synthesis of straight-chain thiophosphoric acid

Figure 10 illustrates synthesis of straight-chain alkylamido-phosphoric

Figure 11 illustrates a synthesis scheme for preparation of

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acid.

conformationally restrained, cyclic phosphate compounds.

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Figure 12 illustrates a synthesis scheme for preparation of conformationally restrained, cyclic phosphate compounds.

Figure 13 illustrates a synthesis scheme for preparation of conformationally restrained, cyclic phosphate compounds. Figure 14 illustrates a synthesis scheme for preparation of conformationally restrained compounds with a free phosphate moiety.

Figure 15 illustrates an alternative synthesis scheme for preparing 2monophosphates. Figure 16 illustrates an alternative synthesis scheme for preparing 1,3-

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Figure 17 illustrates a synthesis scheme for preparing compounds having an —N(H)—acyl group as X3. bisphosphate compounds.

Figure 18 illustrates a synthesis scheme for preparing compounds

Figure 19 illustrates a synthesis scheme for preparing compounds having an —N(H)—imidazole group as X³.

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having an —N(H)—C(O)—O—R7 as X3.

Figure 21 is a graph illustrating the doso-dependent inhibition of LPA-Figure 20 illustrates a synthesis scheme for preparing compounds having an -N(H)-C(S)-O-R7 as X3.

induced chloride currents in Xenopus oocytes by extracellular application of 56 (SAP, 4:0)

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Figure 22 is a graph illustrating the dose-dependent inhibition of LPAinduced chloride currents in Xenopus cocytes by extracellular application of 57 (SAP, 6:81

Figures 23A-B are graphs illustrating the doso-dependent inhibition of LPA-induced chloride currents in Xenopus oocytes by extracellular application of 66 (MAGDP, 18:0). The arrow indicates the time of the intracellular injection of 5 µM 66, followed by the extracellular application of LPA.

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Figure 24 is a graph illustrating dose-inhibitory effect of 66 (MAGDP, increasing amounts of 66. Data points represent the peak amplitude of the measured 18:0). A constant amount of LPA (5 nM) was applied to oocytes together with bloride currents.

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Figure 25 is a graph illustrating the dose-dependent inhibition of LPAinduced chloride currents in Xenopus oocytes by extracellular application of 92 MAGDP, 22:0).

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Figure 26 is a graph illustrating the dose-dependent effect of 56a (SDAP, 14:0/2:0) on Xenopus oocytes.

migration. Test compound concentration was 1 µM; LPA concentration was 0.1 µM. Figure 27 is a bar graph depicting the effects of compounds 56 (SAP, 14:0), 56a (SDAP, 14:0/2:0), and 66 (MAGDP, 18:0) on LPA-induced HEY cell

Ca^{2*} responses in RH7777 cells heterologously expressing Edg-2 (28A), Edg 4 (28B), or Edg -7 (28C). Each data point represents the average of at least three measurements Figures 28A-C are graphs illustrating the dose response relationship for

RH7777 cells, expressing Edg-2, -4, or -7, were exposed to a mixture of 100 nM LPA responses elicited by LPA in Edg-2 and -7, but not Edg-4 expressing RH7777 cells. Representative Ca2 responses are shown for stable Edg-2 (29A), Edg-4 (29B), and Figures 29A-D are graphs illustrating DGPP 8:0 inhibition of Ca2+ 18:1 and 1 µM DGPP 8:0. Control cells were exposed to 100 nM LPA 18:1. Edg-7 (29C) expressing cells, or cells transiently expressing Edg-4 (29D).

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LPA 18:1 mixed with increasing concentrations of DGPP 8:0 and the peak area of the esulting Ca2* responses were measured (30A). Cells were also exposed to increasing characterization of the inhibition of the LPA response by DGPP 8:0 in RH7777 cells 1M concentration of the indicated lipid (30C). The peak areas of the Cath responses concentrations of LPA 18:1 mixed with a 500 nM concentration of DGPP 8:0 (30B). Edg-7 cells were exposed to a 250 nM concentration of LPA 18:1 mixed with a 500 expressing Edg-7 (Edg-7 cells). Cells were exposed to a 250 nM concentration of are represented as the average values of a minimum of three measurements ± SD. Figures 30A-C are graphs which illustrate the pharmacological

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expressing Edg-2 (Edg-2 cells). Stable Edg-2 cells exposed to a 250 nM concentration of LPA 18:1 mixed with increasing concentrations of DGPP 8:0 and peak areas of the haracterization of the inhibition of the LPA response by DGPP 8:0 in RH7777 cells concentration of the indicated lipid (31C). Responses are represented as the average concentrations of LPA 18:1 mixed with a 10 µM concentration of DGPP 8:0 (31B). 2dg-2 cells exposed to a 250 nM concentration of LPA 18:1 mixed with a 10 µM Ca2* responses were measured (31A). Edg-2 cells were exposed to increasing /alues of a minimum of three measurements ± S.D.

Figures 31A-C are graphs which illustrate the pharmacological

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the indicated lipids (32A). Cells transiently expressing Edg-4 cells were exposed to a exposed to a 500 nM concentration of LPA 18:1 mixed with a 5 µM concentration of 100 nM concentration of LPA 18:1 mixed with a 1 µM concentration of the indicated relationship for DGPP in Edg.4-expressing RH7777 cells. Stable Edg.4 cells were Figures 32A-B are graphs which illustrate the structure-activity

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PCT/US01/08729

-16-

PCT/US01/08729

lipids (32B). The peak areas of the Ca^{2*} responses were measured and are represented as the average values of a minimum of three measurements \pm S.D.

Figures 33A-C are graphs which illustrate the pharmacological characterization of DGPP 8:0 on the LPA-elicited CI currents in *Xenopus* oncytes. Occytes were exposed to a 5 nM concentration of LPA 18:1 mixed with increasing concentrations of DGPP 8:0 and the peak amplitude of the resulting oscillatory Ci currents were measured (33A). Occytes were exposed to increasing concentrations of LPA 18:1 mixed with a 200 nM concentration of DGPP 8:0 (33B). Data points represent the average values of a minimum of three measurements ± S.D. Occytes were treated with 5 nM LPA 18:1, or a mixture of 5 nM LPA 18:1 and 1 µM DGPP 8:0 as indicated (33C). The intracellular injection of 1 µM DGPP 8:0 is indicated by the arrow.

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Figures 34A-D are graphs which illustrate DGPP 8:0 inhibiting the LPA-elicited Ca** responses in NIH3T3 fibroblasts and HEY ovarian cancer cells. RT-PCR analysis of NIH3T3 cells for Edg and PSP24 receptor transcripts (34A). NIH3T3 cells were exposed to a 100 nM concentration of LPA 18:1, or S1P, mixed with a 10 µM concentration of DGPP 8:0 (34B). RT-PCR analysis of HEY cells for the presence of the Edg and PSP24 transcripts (34C). HEY cells were exposed to a 100 nM concentration of LPA 18:1, or S1P, mixed with a 1 µM concentration of DGPP 8:0 (34D). The peak areas of the resulting Ca** responses were measured and are represented as the average of a minimum of three measurements ± S.D.

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Figure 35 is a graph illustrating DGPP 8:0 inhibition of LPA-elicited proliferation of NIH3T3 cells. NIH3T3 cells were scrum-starved for 6 hr and exposed to a 5 µM concentration of LPA 18:1 mixed with a 10 µM concentration of the indicated lipids. Control cells received solvent (BSA) in place of LPA 18:1. The cells were incubated for 24 hr with the lipids and counted. Data are representative of three experiments.

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Figure 36 is a graph which illustrates the pharmacological characterizztion of the inhibition of the LPA response by straight-chain fatty acid phosphate compounds 106-110 in Xenopus oocytes.

Figure 37 is a graph which illustrates the pharmacological characterization of the inhibition of the LPA response by straight-chain fatty acid phosphate compound 108 in Xenopus oocytes.

Figure 38 is a graph illustrating the pharmacological characterization of the antagonist or agonist induced response of RH7777 cells inidividually expressing Edg-2, Edg-4, or Edg-7 receptors, following exposure of the cells to straight-chain fatty acid phosphate compound 108. Peak areas of the Ca²⁺ responses were measured.

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WO 01/71022 - 17 -

PCT/US01/08729

DETAILED DESCRIPTION OF THE INVENTION

One aspect of the present invention relates to a compound according to

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wherein,

at least one of
$$X^1$$
, X^2 , and X^3 is $(HO)_2PO-Z^1-$ or $(HO)_2PO-Z^2-P(OH)O-Z^1-$, X^1 and X^2 are linked together as $-O-PO(OH)-$ O-, or X^1 and X^3 are linked together as $-O-PO(OH)-NH-$;

at least one of X¹, X², and X³ is R¹—Y¹—A— with each being the same or different when two of X¹, X², and X³ are R¹—Y¹—A—, or X² and X³ are linked together as —N(H)—C(O)—N(R¹)—;
optionally, one of X¹, X², and X³ is H;

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to 30, or O;

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A is either a direct link, $(CH_2)_k$ with k being an integer from 0

 Y^{l} is $-(CH_{2})$ — with l being an integer from 1 to 30, -0—,

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$$Z^1$$
 is $-(CH_2)_m$ — or $-O(CH_2)_m$ — with m being an integer from 1 to 50, $-C(R^3)H$ —, $-NH$ —, $-O$ —, or $-S$ —;

$$Z^2$$
 is $-(CH_2)_n$ or $-O(CH_2)_n$ with n being an integer from

1 to 50 or —O—;

 Q^1 and Q^2 are independently H_2 , $=NR^4$, =0, a combination of H and $-NR^5R^6$;

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R¹, for each of X¹, X², or X³, is independently hydrogen, a straight or branched-chain Cl to C30 alkyl, a straight or branched-chain C2 to C30 alkenyl, an aromatic or heteroaromatic ring with or without mono-, di-, or trisubstitutions of the ring, an acyl including a Cl to C30 alkyl or an aromatic or heteroaromatic ring, an arylalkyl including straight or branched-chain Cl to C30 alkyl, an aryloxyalkyl including straight or branched-chain Cl to C30 alkyl,

- 18 -

PCT/US01/08729

straight or branched-chain C1 to C30 alkyl groups as described above, with the alkyl Arylalkyl and aryloxyalkyl groups can include, without limitation, group binding to the Y' group of the R'-Y'-A- chain.

compound according to formula (I) are the following previously known endogenous or synthetic compounds: lysophosphatidic acid, phosphatidic acid, cyclic phosphatidic acid, alkenyl glyerolphosphate, dioctyl-glycerol pyrophosphate, and N-palmitoyl-L. Specifically excluded from the above-identified definition of the

Exemplary compounds according to formula (I) are the subclass compounds according to formulae (II)-(V) below.

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one of X', X², and X² is (HO),PO-Z²-P(OH)O-Z¹-, with Z¹ and Z² being O; and two of X¹, X², and X³ are R¹—Y¹—A—, with A being a direct link and Y¹ being O In the structures of formulae (II)A and (II)B, Q1 and Q2 are both H2; for each. Each R1 is defined independently as above for formula (I).

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for formula (I). Preferred species of within the scope of formula Π are where X^3 is lirect link and Y' being -NH- for each. Each R' is defined independently as above (HO),PO-Z1-, with Z1 being O; and X2 and X3 are R1-X1-A-, with A being a NH₂ and X^2 is —NHR¹ with R¹ being a C14 to C18 alkyl, more preferably either a In the structures of formula (III), Q1 is H2; Q2 is =0; X1 is ន

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neteroaromatic ring, an arylalkyl including straight or branched-chain C1 to C30 alkyl, R2, R3, R4, R5, R6, R7, and R8 are independently hydrogen, a straight or branched-chain C1 to C30 alkyl, a straight or branched-chain C2 to C30 alkenyl, an aromatic or heteroaromatic ring with or without mono-, di-, or trisubstitutions of the ring, an acyl including a C1 to C30 alkyl or aromatic or or an aryloxyalkyl including straight or branched-chain C1 to C30 alkyl.

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where xa and xb each are from 0 to 27 and (xa+xb) is not more than 27; and branched except that one or more CH2 groups are replaced by CHW groups where W is an alkyl For each of the above-identified R groups (e.g., R1 - R8), it is intended hat one or more CH2 groups are replaced by CHW groups or a CH group is replaced side chain; straight chain alkenyls have the formula — (CH2), CH=CH(CH2), CH3 chain alkenyls have the formula as defined above for straight chain alkenyl, except that straight chain alkyls have the formula $-(CH_2)_1CH_3$ where x is from 0 to 29; tranched chain alkyls have the formula as defined above for straight chain alkyl, by a CW group, where W is an alkyl side chain.

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aromatic or heteroaromatic rings can include mono-, di-, or tri-substitutions of the ring Aromatic or heteroaromatic rings include, without limitation, phenyls, pyrrolidines, piperidines, thiophenes, furans, napthals, bi-phenyls, and indoles. The located at the ortho, meta, or para positions on the rings relative to where the ring binds to the Y1 group of the R1-Y1-A-chain. Substitutions on the rings can include, without limitation, alkyl, alkoxy, amine (including secondary or tertiary indenes, pyrroles, imidazoles, oxazoles, pyrrazoles, pyridines, mines), alkylamine, amide, alkylamide, acids, alcohols.

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Acyl groups can include either alkyl, alkenyl, or aromatic or heteroaromatic rings as described above.

-20-

PCT/US01/08729

C14 allyl or a C18 allyl; or where X^3 is $-NHR^1$ with R^1 being an acetyl group and X^2 is $-NHR^1$ with R^1 being a C14 allyl.

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In the structures of formula (IV), Q^1 is $=NR^4$, Q^2 is H_2 ; X^1 and X^2 are linked together as -O-PO(OH)-O-; and X^3 is R^1-Y^1-A- , with A being a direct link and Y^1 being -NH-. R^1 and R^4 are as defined above for formula (I).

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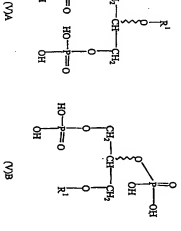
In the structures of formulae (V)A and (V)B, Q^1 and Q^2 are both H_2 : two of X^1 , X^2 , and X^3 are (HO)₂PO— Z^1 —, with Z^1 being O for each; and one of X^1 , X^2 , and X^2 is R^1 — Y^1 —A—, with A being a direct link and Y^1 being —O—. R^1 is as defined above for formula (I). Preferred species within the scope of formulae (V)A and (V)B include the compounds where R^1 is an acyl including a C21 alkyl or where R^1 is a C18 alkyl.

15

WO 01/71022

PCT/US01/08729

-21



The compounds according to formula (I), as well as the subgenus compounds according to formulae (II)A, (II)B, (III), (IV), (V)A, and (V)B, can be prepared using the synthesis schemes described below.

To synthesize the serine amides (SA) and serine amide phosphate (SAP) series (formula (III)), the procursor ι-Boc protected β-lactone (25) was first synthesized. Starting with commercially available ι-Boc-L-serine (Figure 2, 24), triphenyl phosphine (PPh₃) and diethylazidodicarboxylate (DEAD) were introduced under Mitsunobo conditions, affording compound 25 in ca. 50% yield (Sun et al., 1996). Attempts using procedure developed by Sun et al. to open the highly labile β-lactone 25 with various primary amines to obtain hydroxy amides 26-34 failed, in spite of using various reagents (triethyl amine, etc.). Instead, by refluxing the primary amines with the β-lactone in THr, the ι-Boc protected hydroxy amides 26-34 were obtained. Compounds 26-34 were purified using flash column chromatography. Trifluoroacetic acid (TFA)-mediated removal of the ι-Boc protecting group afforded compounds 35-43 as TFA salts.

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To synthesize compounds \$5-59, the t-Boc protected hydroxy amides 26-30 were phosphorylated. A careful study of the final compound suggested that the final compound would possess a highly hydrophobic region and a highly hydrophilic region. Both regions may cause problems during the extraction process and/or attach to the column during the purification stage. To circumvent these potential problems, phosphoramidate chemistry was employed. By using phosphoramidate chemistry, it was hypothesized that the phosphate hydroxyl groups could be protected to render the molecule completely hydrophobic, thereby facilitating its smooth purification.

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WO 01/71022

PCT/US01/08729

Essentially, a combination of procedures was used to obtain the desired products (55-59) (Lynch et al., 1997; Bittman et al., 1996; Liu et al., 1999). Starting hydroxyamides (26-30) were repeatedly washed with anhydrous pyridine, and dried in high vacuum for over 48 hrs. The pyridine-washed hydroxyamides were maintained under an atmosphere of argon. 1H-tetrazole and a freshly distilled 1:1 mixture of THF/CH₂Cl₂ were then added. The phosphorylating agent, dibenzyldiisopropyl phosphoramidate, was added. After monitoring the reaction by TLC, the phosphonate was oxidized to the phosphate in situ with peracetic acid. The reaction mixture was purified via column chromatography to afford compounds 50-54 as benzyl-protected phosphates. The removal of the protecting benzyl groups was carried out in ethanol by subjecting compounds 50-54 to catalytic reduction using 10% palladium on activated carbon (Pd/C) under H₂ atmosphere at 60 psi to yield compounds 55-59 (Figure 3). Reacting 56 with aceric anhydride afforded compound 56a (Figure 3).

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Once the phosphorylation technique was elucidated for the synthesis of the SAP series (compounds \$5-59), a similar procedure was used for the synthesis of bisphosphates (formulae (V)A and (V)B) (Figures 4 and 5A-B). The commercially available diols 60-62 were washed with anhydrous pyridine, and were dried for 48 hrs under high vacuum. These dried diols (60-62) were dissolved in freshly distilled 1:1 THF/CH₂Cl₂, followed by the addition of 1H-terazole. To this stirred mixture was added dibenzy/diisopropyl phosphoramidate. The reaction mixture was monitored via TLC, and at the appropriate time the phosphonate was oxidized to the phosphate in situ with peracetic acid. The reaction mixture was purified with column chromatography to afford compounds 63-65 as benzyl-protected bisphosphates. The removal of the protecting benzyl groups was carried out in ethanol by subjecting compounds 63-65 to catalytic reduction using 10% palladium on activated carbon (Pd/C) under H₂ atmosphere at 60 psi to yield compounds 66-68 was followed to obtain compounds 85-92.

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While compounds 85-92 are 1,2-biphosphates, Figure SB illustrates the synthesis of 1,3-biphosphates. Commercially available 2-phenoxy-1,3-propano-diol was used as the starting material. The starting compound was first protected with *t*-BuOK in the presence of methyl iodide, followed by catalytic hydrogenation to give an intermediate which was then reacted with a halide (RX, where R is as defined above for R¹). The recovered intermediate was subsequently treated with AlCl₃ in the presence of ethyl-SH to yield a 1,3 diol possessing the RO group bound to C2 of the backbone. The recovered 1,3 diol was dissolved in freshly distilled 1:1 THF/CH₂Cl₂, followed by the addition of 1H-tetrazole. To this stirred mixture was added

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dibenzyldiisopropyl phosphoramidate. The reaction mixture was monitored via TLC, and at the appropriate time the phosphonate was oxidized to the phosphate in situ with peracetic acid. The reaction mixture was purified with column chromatography to afford benzyl-protected bisphosphate compounds. Removal of the protecting benzyl groups was carried out in chanol by subjecting the compounds to catalytic reduction using 10% palladium on activated carbon (Pd/C) under H₂ atmosphere at 60 psi to yield 1,3-bisphosphate compounds.

To synthesize the pyrophosphates of formulae (II)A and (II)B, glycidal tosylate ((2R)(-) or (2R)(+)) was used as the starting material (Figures 6A-B). Opening of the ring was catalyzed by a Lewis acid, such as BF3, in the presence of an alcohol, affording an intermediate which was tosylate-protected at the C1 position. In the next step, the alcohol at the C2 position was replaced with an R group (e.g., R¹ as described above) using as excess of R-triflate and 2,6-di-tert-butyl-4-methylpyridine, affording the di-ether intermediate. Treament of the di-ether intermediate with tris(tetran-butylammonium) bydrogen pyrophosphate caused nucleophilic attack of the tosylate, replacing the tosylate with a pyrophosphate substituent at the C1 position.

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To produce the pyrophosphate of formula (II)B, the tosylate protected intermediate was treated with benzyl alcohol in the presence of triflic anhydride and 2,6-di-tert-butyl-4-methylpyridine, which benzylates the intermediate at the C2 position. The tosylate protecting group on the benzylate intermediate was removed first by the action of potassium superoxide in the presence of 18-crown-6, affording a hydroxyl group at the C1 position which was subject to replacement with an R group (e.g., R¹ as described above) using an excess of R-triflate and 2,6-di-tert-butyl-4-methylpyridine. The resulting di-ether intermediate still possessed the benzyl protecting group at the C2 position. The benzyl motecting group was removed by

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incur) pyramic. The resulting ur-ener internetiate stut possessed are penzyl protecting group at the C2 position. The benzyl protecting group was removed by hydrogenation and the subsequent hydroxyl group was tosylated by the action of pyridine and p-toluenesulfonyl chloride, producing a di-ether bearing a tosyl group at the C2 position. The tosylate group was removed by nucleophilic attack upon treatment with tris(tetra-n-butylanmonium) hydrogen pyrophosphate, replacing the tosylate with a pyrophosphate substituent at the C2 position.

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Alternative schemes for preparing phosphates and biphosphates (as well as pyrophosphates, phosphonates, etc.) are illustrated in Figures 15 and 16.

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In Figure 15, glycidal bromide was used as the starting material along with an alcohol (ROH). The reaction conditions included treatment with K₂CO₃ followed by treatment with the ammonium salt C₆H₆CH₂N'(C₂H₃)₂CI; resulting in displacement of the bromide with the R group. The ring of the glycidal intermediate was then opened following treatment with 1M HCl in ether and an alcohol (R¹OH),

PCT/US01/08729

psi to yield monophosphate compounds. carried out in ethanol by subjecting the benzyl-protected phosphates to catalytic afford benzyl-protected phosphates. The removal of the protecting benzyl groups was di-ether was mixed with 1H-tetrazole and to this stirred mixture was added reduction using 10% palladium on activated carbon (Pd/C) under $m H_2$ atmosphere at 60 peracetic acid. The reaction mixture was purified with column chromatography to and at the appropriate time the phosphonate was oxidized to the phosphate in situ with dibenzyldiisopropyl phosphoramidate. The reaction mixture was monitored via TLC which afforded a di-ether intermediate having a hydroxy group at the C2 postion. The

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of reacting the glycidal bromide with an alcohol (ROH); BnOH was used to protect the psi to yield 1,3 bisphosphates. reduction using 10% palladium on activated carbon (Pd/C) under $m H_2$ atmosphere at 60 was carried out in ethanol by subjecting the benzyl-protected phosphates to catalytic affords benzyl-protected phosphates. The removal of the protecting benzyl groups peracetic acid. The reaction mixture was purified with column chromatography to and at the appropriate time the phosphonate was oxidized to the phosphate in situ with dibenzyldiisopropyl phosphoramidate. The reaction mixture was monitored via TLC was combined with 1H-tetrazole and to this stirred mixture was added activated carbon (Pd/C) under H_2 atmosphere at 60 psi to yield a 1,3 diol. The diol intermediate was de-protected via catalytic reduction using 10% palladium on intermediate having an R group attached via ether bond at the C2 position. This The di-ether was mixed with a halide salt (RX) in aqueous K2CO3, yielding a protected C1 site. The resulting di-ether intermediate has a hydroxy group at the C2 postion. following treatment with 1M HCl in ether and annhydrous BnOH, which protected the bromide with the Bu group. The ring of the glycidal intermediate was then opened with the ammonium salt CoHoCHoN (CoHo)CI; resulting in displacement of the C3 site. The reaction conditions included treatment with K2CO3 followed by treatment In Figure 16, a similar reaction scheme was employed, except instead

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can be attached at the C1 site upon removal of the tosyl group. As shown in Figure bearing ${\bf R}$ and ${\bf R}^1$ substituents), a number of modified phosphates and phosphonates group at the C1 site. As shown in Figure 7B, the intermediate is reacted under basic are removed following treatment with TMSBr, affording a $-(R^3)CH-PO(OH)_2$ PO(0—protecting group)2 to form a single bond to the C1 site. The protecting groups remove the tosylate protecting group and allow the modified phosphate $-Z^1$ protecting grouph where Z^l is $-(R^l)CH$ —and X^e is H. The basic conditions 7A, the intermediate is reacted under basic conditions with X⁴-Z¹-PO(0-Using the di-ether intermediate prepared as shown in Figure 6A (e.g.

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where Z^1 is --O, Z^2 is $--CH_2$, and X^4 is H. The basic conditions remove the conditions using tris(tetra-n-butylammonium) with $X^4-Z^1-PO(OH)-Z^2-PO(OH)$, ĸ

tosylate protecting group and allow the modified phosphonate

and water wash, affording a -OCH2CH2-PO(OH)2 group at the C1 site. modified phosphate —Z¹—PO(O—protecting group); to form a single bond to the CI X^{\bullet} is H. The basic conditions remove the tosylate protecting group and allow the conditions with X⁴-Z¹-PO(O-protecting group), where Z¹ is -OCH₂CH₂- and site. The protecting groups are removed following treatment with TMSBr in collidine installed at the C1 site. As shown in Figure 7C, the intermediate is reacted under basic with acidic conditions and CH3CN, the -O-PO(OH)-CH2-PO(OH), group is $-Z'-PO(OH)-Z'-PO(OH)_2$ to form a single bond to the C1 site. Upon treatment

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the resulting product was treated with di-tert-butyl diisopropylphosphoramidate, scheme illustrated in Figure 11. Compounds 26-30 were reacted with 1H-tetrazole and of formula (III), compounds 26-30 were used as starting materials in the synthesis the compounds of formula (III). peracetic acid yielded a cyclic phosphate intermediate. Reduction with TFA yielded causing an intramolecular cyclization. In situ oxidation of the phosphonate with To prepare the conformationally restricted cyclic-phosphate compound

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hydroxyl group bound to the C2 carbon. The cyclic intermediate is then treated with cyclic phosphates where X¹ and X² together are --O-PO(OH)--O-. A benzylunder H_2 atmosphere (as described above) affords a cyclic phosphate bearing a intramolecular cyclization. Treatment with 10% palladium on activated carbon (Pd/C) protected 1,3 diol intermediate is reacted with POCl3, which results in an Other conformationally restricted compounds can also be prepared. As shown in Figure 12, an alternative scheme is shown for preparing

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an excess of R-triflate and 2,6-di-tert-butyl-4-methylpyridine to afford the final

tris(1,2,4,-triazole)phosphate followed by 2% HCl wash, resulting in intramolecular phosphate where X^1 and X^3 together are -0—PO(OH)—NH—. Using the intermediates 35-43 prepared above as starting material, they are treated with As shown in Figure 13, a scheme is shown for preparing a cyclic

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as starting materials, they are treated with anhydrous COCl2, which inserts a carbonyl together are --N(H)--C(O)-N(R1)-. Using the intermediates 50-54 prepared above compound where the phosphate group is not a part of the ring; specifically, X^2 and X^3 between between the amines bound to the C2 and C3 carbons during cyclization. As shown in Figure 14, a scheme is shown for preparing a cyclic

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WO 01/71022

27

PCT/US01/08729

Benzyl protecting groups are removed from the phosphate using 10% palladium on activated carbon (Pd/C) under H_2 atmosphere (as described above).

Another class of compounds which can be used as agonists or antagonists of the LPA receptors are fatty acid phosphates or straight-chain phosphates. As shown in Figure 8, anhydrous n-alkanol and 1*H*-tetrazole can be dissolved in anhydrous methylene chloride. A solution of dibenzyl-N,N-diisopropyl phosphoramidite in anhydrous methylene chloride can be added. Subsequently, peracetic acid in anhydrous methylene chloride can be added dropwise to afford the benzyl-protected fatty acid phosphates 101-105. The benzyl-protecting groups are removed following treatment in anhydrous methanol with 10% palladium on activated carbon (Pd/C) under H₂ atmosphere (as described above), affording the fatty acid phosphates 106-110.

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enzyl-protecting groups are removed following treatment in anhydrous methanol with can be added dropwise to afford the benzyl-protected fatty acid amidophosphates. The senzyl-protecting groups are removed following treatment in anhydrous methanol with As an alternative to preparing fatty acid phosphates, thiophosphates and .0% palladium on activated carbon (Pd/C) under H2 atmosphere (as described above), abloride can be added. Subsequently, peracetic acid in anhydrous methylene chloride chloride can be added. Subsequently, peracetic acid in anhydrous methylene chloride nercaptoalkanes and 1H-tetrazole can be dissolved in anhydrous methylene chloride. 10% palladium on activated carbon (Pd/C) under H2 atmosphere (as described above), can be added dropwise to afford the benzyl-protected fatty acid thiophosphates. The iffording the fatty acid thiophosphates. As shown in Figure 10, for example, an nilkylamine and 1H tetrazole can be dissolved in anhydrous methylene chloride. A A solution of dibenzyl-N,N-diisopropyl phosphoramidite in anhydrous methylene solution of dibenzyl-N,N-diisopropyl phosphoramidite in anhydrous methylene amidophosphates can also be prepared. As shown in Figure 9, for example, nffording the fatty acid amidophosphates.

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Bach of the above-identified reaction schemes can be further modified by attacking a primary amine group as shown in Figures 17-20. The intermediate is prepared, e.g., from compounds 50-54 which were treated with TFA to remove the t-Boc protecting group, affording the primary amine at the C2 site while leaving the phosphate protected.

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In Figure 17, the intermediate compound possessing a primery amine at the C2 position is attacked with an acid halide (e.g., R¹COCI), which converts the primary amine into an amide (—N(H)—C(O)—R¹). The benzyl-protected phosphate

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can then be de-protected using treatment with 10% palladium on activated carbon (Pd/C) under H_2 atmosphere (as described above).

In Figure 18, the intermediate compound possessing a primery amine at the C2 position is attacked with N-acetyl imidazoline in POCl3, which converts the primary amine into a secondary amine (—N(H)—imidazole). Substituted imidazolines can also be used. The benzyl-protected phosphate can then be deprotected using treatment with 10% palladium on activated carbon (Pd/C) under H2 atmosphere (as described above).

In Figure 19, the intermediate compound possessing a primery amine at the C2 position is attacked with R¹OC(O)Cl, which converts the primary amine into an carbamate (—N(H)—C(O)—O—R¹). The benzyl-protected phosphate can then be deprotected using treatment with 10% palladium on activated carbon (Pd/C) under H₂ atmosphere (as described above).

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In Figure 20, the intermediate compound possessing a primery amine at the C2 position is attacked with R¹NCO or R¹NCS, which converts the primary amine into either a uramide (—N(H)—C(O)—N(H)—R¹) or thiouramide (—N(H)—C(S)—N(H)—R¹). The benzyl-protected phosphate can then be de-protected using treatment with 10% palladium on activated carbon (Pd/C) under H₂ atmosphere (as described above).

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Thus, the non-cyclic compounds of the present invention can be prepared by reacting (Y²O)₂PO—Z¹¹¹ or (Y²O)₂PO—Z¹² —P(OH)O—Z¹¹ = Z¹¹ where Z¹¹ is —(CH₂)_m— or —O(CH₂)_m— with m being an integer from 1 to 50, — C(R³³)H—, or —O—, Z¹² is —(CH₂)_m— or —O(CH₂)_m— with n being an integer from 1 to 50 or —O—, Z¹² is H or a first leaving group or —Z¹¹—Z¹¹ together form the first leaving group, and Y² is H or a protecting group; with an intermediate compound according to formula (VI), followed by a de-protection step, if necessary, both performed under conditions effective to afford a compound according to formula (I) where one or two of X¹, X², and X² is (HO)₂PO—Z¹— or (HO)₂PO—Z²— P(OH)O—Z¹— where Z¹ and Z² being defined as above.

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The intermediate compound of formula (VI) has the following structure:

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 X^{12} and X^{13} are linked together as $-N(H)-C(O)-N(R^{11})-$; being the same or different when two of X11, X12, and X13 are R11-Y11-A-, or at least one of X11, X12, and X13 is R11-Y11-A- with each

leaving group; at least one of X11, X12, and X13 is OH, NH2, SH, or a second

optionally, one of X11, X12, and X13 is H;

A is either a direct link, (CH2)k with k being an integer from 0

 Y^{11} is $-(CH_2)$ — with l being an integer from 1 to 30, -0—,

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to 30, or O;

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H and -NR 14 R 15;

Q1 and Q2 are independently H2 =NR13, =O, a combination of

substitutions of the ring, an acyl including a C1 to C30 alkyl or an aromatic or alkenyl, an aromatic or heteroaromatic ring with or without mono-, di-, or triheteroaromatic ring, an arylalkyl including straight or branched-chain C1 to C30 alkyl, straight or branched-chain C1 to C30 alkyl, a straight or branched-chain C2 to C30 \mathbb{R}^{11} , for each of \mathbb{X}^{11} , \mathbb{X}^{12} , or \mathbb{X}^{13} , is independently hydrogen, a

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an aryloxyalkyl including straight or branched-chain C1 to C30 alkyl.

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arylalkyl including straight or branched-chain C1 to C30 alkyl, or an aryloxyalkyl aromatic or heteroaromatic ring with or without mono-, di-, or tri-substitutions of the branched-chain C1 to C30 alkyl, a straight or branched-chain C2 to C30 alkenyl, an ring, an acyl including a C1 to C30 alkyl or aromatic or heteroaromatic ring, an R12, R13, R14, R15, R16, and R17 are independently hydrogen, a straight or 置

including straight or branched-chain C1 to C30 alkyl.

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WO 01/71022

PCT/US01/08729

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compound(s), together with the carrier, excipient, stabilizer, etc. stabilizers, and can be in solid or liquid form such as, tablets, capsules, powders, invention. The pharmaceutical composition can also include suitable excipients, or includes a pharmaceutically-acceptable carrier and a compound of the present present invention, such compounds can be used to prepare pharmaceutical about 0.01 to 99 percent, preferably from about 20 to 75 percent of active solutions, suspensions, or emulsions. Typically, the composition will contain from further aspect of the present invention relates to a pharmaceutical composition that compositions suitable for treatment of patients as described hereinafter. Therefore, a Having prepared the LPA receptor agonists and antagonists of the

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magnesium stearate. cornstarch, potato starch, or alginic acid, and a lubricant, like stearic acid or with binders like acacia, comstarch, or gelatin, disintegrating agents, such as with conventional tablet bases such as lactose, sucrose, or comstarch in combination the present invention and a carrier, for example, lubricants and inert fillers such as, form can be a capsule, such as an ordinary gelatin type containing the compounds of lactose, sucrose, or cornstarch. In another embodiment, these compounds are tableted The solid unit dosage forms can be of the conventional type. The solid

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vegetable, or synthetic origin, for example, peanut oil, soybean oil, or mineral oil. In for injectable solutions. as propylene glycol or polyethylene glycol, are preferred liquid carriers, particularly surfactant and other pharmaceutically and physiologically acceptable carrier, including a physiologically acceptable diluent with a pharmaceutical carrier. Such carriers injectable or topically-applied dosages by solution or suspension of these materials in general, water, saline, aqueous dextrose and related sugar solution, and glycols, such adjuvants, excipients or stabilizers. Illustrative oils are those of petroleum, animal, include sterile liquids, such as water and oils, with or without the addition of a The compounds of the present invention may also be administered in

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may be administered in a non-pressurized form such as in a nebulizer or atomizer. or suspension may be packaged in a pressurized aerosol container together with suitable propellants, for example, hydrocarbon propellants like propane, butane, or isobutane with conventional adjuvants. The materials of the present invention also For use as aerosols, the compounds of the present invention in solution

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instillation, by intracavitary or intravesical instillation, intraocularly, intraarterially, subcutaneously, intravenously, intramuscularly, intraperitoneally, by intranasal present invention can be administered orally, topically, transdermally, parenterally, Depending upon the treatment being effected, the compounds of the

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WO 01/71022 - 30 - CT/US01/08729

intralesionally, or by application to mucous membranes, such as, that of the nose, throat, and bronchial tubes.

Compositions within the scope of this invention include all compositions wherein the compound of the present invention is contained in an amount effective to achieve its intended purpose. While individual needs vary, determination of optimal ranges of effective amounts of each component is within the skill of the art. Typical dosages comprise about 0.01 to about 100 mg/kg-body wr. The most preferred dosages comprise about 0.1 to about 100 mg/kg-body wr. The most preferred dosages comprise about 1 to about 100 mg/kg-body wr. Treatment regimen for the administration of the compounds of the present invention can also be determined readily by those with ordinary skill in art.

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Certain compounds of the present invention have been found to be useful as agonists of LPA receptors while other compounds of the present invention have been found useful as antagonists of LPA receptors. Due to their differences in activity, the various compounds find different uses. The preferred animal subject of the present invention is a mammal, i.e., an individual belonging to the class Mammalia. The invention is particularly useful in the treatment of human subjects.

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One aspect of the present invention relates to a method of modulating LPA receptor activity which includes providing a compound of the present invention which has activity as either an LPA receptor agonist or an LPA receptor antagonist and contacting an LPA receptor with the compound under conditions effective to modulate the activity of the LPA receptor.

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The LPA receptor is present on a cell which either normally expresses the LPA receptor or has otherwise been transformed to express a particular LPA receptor. Suitable LPA receptors include, without limitation, EDG-2, EDG-4, EDG-7, and PSP-24 receptors. The tissues which contain cells that normally express these receptors are indicated in Table 1 above. When contacting a cell with the LPA receptor agonist or LPA receptor antagonist of the present invention, the contacting can be carried out while the cell resides in vitro or in vivo.

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To heterologously express these receptors in host cells which do not normally express them, a nucleic acid molecule encoding one or more of such receptors can be inserted in sense orientation into an expression vector which includes appropriate transcription and translations regulatory regions (i.e., promoter and transcription termination signals) and then host cells can be transformed with the expression vector. The expression vector may integrate in the cellular genome or simply be present as extrachronosomal nuclear material. Expression can be either

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WO 01/71022 - 31 -

PCT/US01/08729

constitutive or inducible, although constitutive expression is suitable for most purposes.

The nucleotide and amino acid sequences for EDG-2 is known and reported in An et al. (1997b) and Genbank Accession No. U80811, which is hereby incorporated by reference. An EDG-2 encoding nucleic acid molecule has a nucleotide sequence according to SEQ. ID. No. 1 as follows:

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60 120 240 300	3 4 4 8 8 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9	660 720 780 840 900	960 1020 1080 1095
cocagttcan agocatgant atmaccgang tggamagent gaottggmat cautgtttgt totatgtcam cogcogetto cagacttott tgctgggttg	tgttagcaca caacttactg acggatgago tatgggtgct catgggcoccc	otttgtggta gagaatgtot gaagactgtg gttacttota totcottget	aatgagogoo cacagaaago cagcaatgao
cocagitica agocatgaat ataacogaag iggaaagcat gactiggaat cacigitigi totatgicaa cogcogotio cagacitoti igcigggitg	ggagactgac catctgtggc agctccacac tggocatcgt attgttccaa	acttggtgac otttgtggta agaggactat gagaatgtct tgagtcttct gaagactgtg gattggtttt gttacttcta aattcttcct tctcottgct	tactcctacc gcgacaaaga antgagcgcc agtgagacc ccaccggccc cacagaaagc accatctgg ctggagttca cagcaatgac
atggetgeda tototactto catecotgta attteacago gaacacagt gettetacaa egagecatt geettettt ritgeacag aatggaacac agteageago tegtgatgg Atctteatca tgftggeaa cotattggte atggtggea cattteceta tttattacet aatggetaat otggetgera	cccaatactc agcctgacgg ttccgcatgc atctggacta gatattgaan	gccattttca tatgttcgcc gataccatga tggactcctg gcctatgaga	tactcctadd agtgagaacd accatcttgg
tototactto catocotgta gcttctacaa egagtccatt aatggaacac agtcagcaag tgttggccaa cctattggtc ttattacct aatggctaat	caacacagga cattgacacc cattaoggtt cattgtggtc ctgtatctgt	actottactt agtottotgg totatggctca catctttggc ctggaccccg gcggaatcgg ttggggctt tatcatctgc gtccacagtg cgacgtgctg	ctgccatgaa ccccatcatt agatcctetg ctgccagogo oggettcetc cctcaaocac ttbag
tototactto gcttctacaa aatggaacac tgttggccaa tttattacct	atctcatgtt gtcagggcct tcgagaggca tagtggtggt tgggctggaa	actettaett tetatgetea etggaceeg ttggggeett gtecacagtg	ctgccatgaa agatcctctg cggcttcctc tttag
atggotgoca gaaccacagt cttgccacag atcttcatca catttccta	gectaettet tggeteetge getattgea aaceggeggg ataeceagtg	ctctacagtg atggtggttc cggcatagtt gtcattgtgc gacgtgtgct	gaattcaact acctttaggc tcagaccgct cactotgtgg
10	. \$1	70	25

The encoded EDG-2 receptor has an amino acid sequence according to SEQ. ID. No. 2 as follows:

IVC 60			LLA 300		364	
LVMGLGI7	PNTRRLT	YVRORITMS	AYERPELLLA	TILAGVH		
LATEWATVEK	AYPYLMPNTO	MVLYAKIFG	DVCCPQCDVL	SDRBASSIAN		
MALISTSIPV ISODOFTAMN REQCPYNESI AFFYNRSGKH LATBHNIVGK LVMGLGIITVC	PIMLANILU MVAIYVNRRF HPPIYYLMAN LAAADBPAGL AYFYLMFNTG PNTRRLTV6T	MILKUGALLI BIJABVANILI AIALEKHIIV KWOLHIRMB NEKVVVIVV INTMALVRGA IPSVGRNYCIC DIENCENMAD LYSDSYLVIW AIRKIVITVV MVVLYAHIRG YVRORINGMS	RHSSGPRRNR DIMMSLLKIV VIVLARFIIC WIPGLVLLLL DVCCPGCDVL	BFNSAMNPII YSYRDKEMSA TFRQILCCQR SENPTGPTBS SDRSASSLAN TILAGVHSND		
RPQCFYNESI	HPPIYYLAM	LYSDSYLVPW	VIVLGAPIIC	TFRQILCCOR		
ISOPOPTAMA	MVALYVNERP	DIENCENMAP	DIMMSLLKTV	YBYRDKEMSA		
MAISTSIPV	IPIMLANLLV	IPSVGWNCIC	RHSSGPRRNR	RENBAMINEII	HSW	
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The nucleotide and amino acid sequences for EDG4 is known and reported in An et al. (1998b) and Genbank Accession No. NM_004720, which is hereby incorporated by reference. An EDG4 encoding nucleic acid molecule has a nucleotide sequence according to SEQ. ID. No. 3 as follows:

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taacaacagt aotggggctg cgcctccaac tgacctcttc cgactttca	200000000000000000000000000000000000000
alegyteatea tggycaagtg etaetaeaa gagaceateg getteiteta taacaaeagt gygaaagage teageteeca etggygygeo aagagatgyg tgytgygygo abtgyyyetg accyclagog tegtygyet getgaceaat etgetygytea tagaageat gyeteceaac ageogettee accageceat etaetaecetg eteggeaate tggeogogye tgaeetette gygygygygyg octaactet ecteatytee acaategyte eccgateste	
gagaccatog aaggatgtgg ctgctggtca ctcggcatc	The same of the sa
ctactacaac ctggeggcc gctgaccaat ctactacctg cctcatgttc	Canada Contract
toggecoagig teageteea toetggtget accagecat cetacetett	2000
atggtcatca ggcaaagago accgtcageg cgccgcttcc gcgggcgttg	10000

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WO 01/71022 - 32 -

acaotyctgg coatogocyt ggagogycao ogcastytya tygocytyca gctgoacagc 420 cygotycac gruycogycy gytaatyrt attyrgygys tyftygytygy 480 cygotyguty Gyockygaa tototygyaa tygocytyy coctygacog dysteacyc attyrgygysty Gyockygaco troctygyaa tygocyty coctygacog dysteacyc A100 attyrcacoc tygotcacyc cactodygyaa tydocyty coctygacog dysteacyc A100 attyrcacoc tygotcacyc cactodygaa trattictict acygycyga cocyctyty 600 troctyctca tygygygyty gracacocy cattrictict acygygygy gogatygaa 600 cygatycay agoatycay gagacacoc ogcacocyc tygocytyta 780 cygatygra gacacocyc togocacyc togocyctyt 780 cygatycya gagacacyg cagytycyt 780 cygatycya gagatycat gygygytic gyagatyco tygocytyaga aasytactc 900 gagatygog gagacocyc gyacacocyc gyacacocyc gyagatyco tygagatygy cogagytygy cogagytyco cygatycot gyacacocyc gyagatyco gagatycyc cygatycot gyacacocyc gyagatyco gagatycyc cygatycyc attratyc cygagytyc 100 gagatygyoc gacacttco cycyccacyc gyagatycoc ycacacocyc gagaacocyc gagaacocyc gyagatyco accaactyat gyactcocyc gyagatycoc gagaacygy caccacyc yccacocyc 1020 gagatygyc accaacacyc gyactcocyc accaacyg gyactcocyc cattag

The encoded EDG-4 receptor has an amino acid sequence according to SEQ. ID. No. 4 as follows:

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MVIMGQCYYN BTIGDFYNNS GEELGSHWRD EDVDVVALGL TUSULVILIU LLUTALIASN 60
RREEGDYYYL LGELAAADLE AGVAYLELME STOPRTARLS LEGWFLEGGL LDTSLTASVA
TLLALAURH RSYMAVQLES RLEBGRYUML IYGVWYALGL LGLLEBRISHE CLCALDROSR 180
MADLLSRSYL AVMALSSILV PLAWAVYTR IFFYVBREVQ BARESVESTP RYRETTISLV 240
KTYVILIGAS VUCWTJGGOV LLIGGLGCES CSVLAVEKYF LLIABRASIV KRAVYSCEDA 301
EMBRITERELL CCACLROSTR ESVEYTSSAQ GGASTELMLD ENGEPLADST L 351

The nucleotide and amino acid sequences for EDG-7 is known and reported in Bandoh et al. (1999) and Genbank Accession No. NM_012152, which is hereby incorporated by reference. An EDG-7 encoding nucleic acid molecule has a nucleotide sequence according to SEQ. ID. No. 5 as follows:

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The encoded EDG-7 receptor has an amino acid sequence according to SEQ. ID. No. 6 as follows:

PHEPPYYLLA KLAAADEPAG IAYVELMENT GEVSKTILTVN RNELRQGILD SSUTASUTNI. 120

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WO 01/71022 - 33 -

PCT/US01/08729

LVIAVERENG INGHRVESKI TRKKVTLLIL LVWALALENG AVFILGKNCL CHISACSSLA PICSESYLVF-WYVSKLAGEL INVVVYLAIY VYVERKUNUL SPETSGSISR BETENGLKKT VMIVLGAFVV.CWIFGLVVLL ILGIANCROCG VQBYKRWELL LALLMSVVNF IIVSYKDEDM YGYMKUNICC FSQENPERRE SRIESTVLSR SDYGSQYIED SISQGAVCNK STS

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The nucleotide and amino acid sequences for PSP-24 is known and reported in Kawasawa et al. (2000) and Genbank Accession No. AB030566, which is hereby incorporated by reference. A PSP-24 encoding nucleic acid molecule has a nucleotide sequence according to SEQ. ID. No. 7 as follows:

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atgytettet eggentigt gantigegite catacoggya catecaaca aacattigte giglatgaaa acactacat gantigegite catacoggya catecaaca aacattigte giglatgaaa acactacat gantigegite catacoggya catecaaca acgigetegac 120 agtecaatyo tragatalag tritgaacaca gegentika aggetegaa categacate 240 grigatalagta cagetegage cacacaaca gegentika aggetetaa categactet 240 grigatalagta cagetegage categateate attetytite typeatitet 240 grigatalagta categateta categatett 240 grigatalagta categageta acatgatalate attetytitet typeatitet 240 grigatalagta categageta categageta categageta categageta gantitetygg anattettet grigataga aggataggo alcotygiteta taggagatage alcologite aggatalagta categageta categageta gantitetygg agatalagta trecatata gagetalagta salcologite salcologite trecatatagy cancettecate tregategage attetatag categageta trecatate categageta gantiteta cagagageta categageta cagagageta cagagatalag categageta cagagageta tregatete cagagageta trecatate cateacateta cateacatet garagagat cagagatalaga agacattet cagagagacate treatacagaga categageta cacatatett gatecatett 900 grintigagaga treatagag gacatalaga cacatatitet gatecatett 900 aagagaacatet acasaca cacatatitet gatecatett 900 aagacattet acasaca cacatatitet gatecatett 900 aagacattet acacatatotta gantigaga aagacattet acacatatotta gantigaga 1140 aagacateta acacatatotta gantigaga 1140 aagacategaga acatgatega caagacateta acacatatitet gatecatett 900 aagacattet acacatatatet gatecatett 900 aagacattet acacatatotta garacatett 900 aagacattet acacatatotta garacatetti 900 aagacattet acacatatotta garacatetti 900 aagacattet acacatatotta garacatetti 900 aagacattet acacatatotta garacatetat 900 aagacattet acacatatotta garacatetat 900 aagacattet acacatatotta garacatetat 900 aagacatteta acacatatotta 900 aagacatteta acacatatotta 900 aagacatteta acacatatota 900 aagacateta 900 aagacatta 900 aagacatta

The encoded PSP-24 receptor has an amino acid sequence according to SEQ. ID. No. 8 as follows:

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MYSAVLTAF HIGISNITSV VYRNIYANIT LEPDPPQHEDL SPLLRUSSET MAPIGLSSLT 60

VNSTAVETTE ALFKELNIEL QUTLBRIKLE ILEVSSLGRI VYCLMYVOKA AMBRANILL 120

ASLAFADMIL AVLAMPRALV TILITERNIES KSECRVSAMF FWLEVIEGVA ILLIISIDEF 180

LIIVQRQDKL HYPARAVULA VSKATSSCYA FELAVUSEDL QUESBADÇCY POTTYDGYG 240

AVVILSELS SFIPTILLIE AVBIVCHAFF TITSLVATES KEFYQDRFF EISTWLWIG. 360

QMSIDMGEKT, RAFITILLIE AVBIVCHAFF TITSLVATES KEFYQDRFF EISTWLWIG. 360

YLKSALMPLI YYMRIKKUED ACLDMOFRSF KFLEQLFGHT KREINPSAVY VCGHERTVV 419

LPA receptor agonists will characteristically induce LPA-like activity from an LPA receptor, which can be measured either chemically, e.g., Ca²⁺ or Cl current in occytes, or by examining changes in cell morphology, mobility, proliferation, etc. In contrast, LPA receptor antagonists will characteristically block LPA-like activity from an LPA receptor. This too can be measured either chemically, e.g., Ca²⁺ or Cl current in occytes, or by examining changes in cell morphology, mobility, proliferation, etc.

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an LPA receptor with the compound under conditions effective to inhibit LPA-induced induced activity on an LPA receptor. This method includes providing a compound of the present invention which has activity as an LPA receptor antagonist and contacting eceptor antagonists, the present invention also relates to a method of inhibiting LPA. neterologously expresses the receptor. The contacting of the LPA receptor with the activity of the LPA receptor. The LPA recepter can be as defined above. The LPA By virtue of the compounds of the present invention acting as LPA compound of the present invention can be performed either in vitro or in vivo. receptor is present on a cell which normally expresses the receptor or which

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those LPA receptors described above. Therefore, it is expected that the compounds of different cellular pathways which involve signaling through LPA receptors, including As noted above, LPA is a signaling molecule involved in a number of he present invention will modulate the effects of LPA on cellular behavior, either by acting as LPA receptor antagonists or LPA receptor agonists.

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present invention includes those cancers characterized by cancer cells whose behavior receptors. Exemplary forms of cancer include, without limitation, prostate cancer and administering an effective amount of the compound to a patient in a manner effective to treat cancer. The types of cancer which can be treated with the compounds of the is attributable at least in part to LPA-mediated activity. Typically, these types of cancer are characterized by cancer cells which express one or more types of LPA One aspect of the present invention relates to a method of treating cancer which includes providing a compound of the present invention and ovarian cancer

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The compounds of the present invention which are particularly useful for cancer treatment are the LPA receptor antagonists.

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bound by theory, it is believed that the LPA receptor antagonists, upon binding to LPA specific site where cancer cells are present. Thus, administering can be accomplished When administering the compounds of the present invention, they can compounds of the present invention were cytotoxic to prostate cancer cell lines which be administered systemically or, alternatively, they can be administered directly to a in any manner effective for delivering the compound to cancer cells. Without being destroy those cancer cells. As shown in Example 12 infra, several LPA antagonist receptors, will inhibit proliferation or metastasis of the cancer cells or otherwise express one or more LPA receptors of the type described above.

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When the LPA antagonist compounds or pharmaceutical compositions composition can also contain, or can be administered in conjunction with, other of the present invention are administered to treat cancer, the pharmaceutical

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therapeutic agents or treatment regimen presently known or hereafter developed for the treatment of various types of cancer

cells or cell clusters detach from the primary tumor and reach the systemic circulation process, tumor cells must arrest in capillaries, extravasate, and migrate into the stroma to the basement membrane glycoprotein laminin via the cell surface laminin receptors. of the tissue to make secondary foci. First, tumor cells must recognize signals on the endothelial cell that arrest them from the circulation. Second, tumor cells must attach Following attachment to the basement membrane, tumor cells secrete proteases to Cancer invasion is a complex multistep process in which individual or the lymphatics to spread to different organs (Liotta et al., 1987). During this

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correlation (Hoffman-Wellenhof et al., 1995). It is a well-documented fact that PLGFs component that is fully capable of restoring tumor cell invasion in serum-free systems promote proliferation and increase invasiveness of cancer cell in vitro. Imamura and tumor cell invasion and metastasis. The relationship between motility of tumor cells in vitro and the metastatic behavior in animal experiments indicates a strong direct degrade the basement membrane. Following attachment and local proteolysis, the third step of invasion is tumor cell migration. Cell motility plays a central role in colleagues established that cancer cells require serum factors for their invasion (Imamura et al., 1991), and later identified LPA as the most important serum (Xu et al., 1995a; Imamura et al., 1993; Mukai et al., 1993).

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presence of EDG-2 and EDG-7 receptors in these cell lines. Recently, Im et al. (2000) demonstrated that BDG-7 is expressed in prostate cancer cell lines; namely, PC-3 and LNCaP cells. RT-PCR analysis on the prostate cancer cell lines DU-145, PC-3, and cancer cell lines, whereas EDG-3 is present in LNCaP and DU-145 prostate cancer It has been shown that PLGFR are expressed in ovarian cancer cell LNCaP lines showed that EDG-2, 4, 5, and EDG-7 are present in all three prostate ines; namely, OCC1 and HEY cells. Specifically, RT-PCR analyses show the cell lines.

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specific ovarian cancer cell lines. Thus, the LPA antagonists of the present invention As shown in the Examples, several LPA receptor antagonists of the novide an alternative approach for treatment of LPA-mediated cancers, including present invention are capable of targeting specific prostate cancer cell lines and prostate cancer and ovarian cancer.

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mbancing cell proliferation. This method of enhancing cell proliferation includes the steps of providing a compound of the present invention which has activity as an agonist of an LPA receptor and contacting the LPA receptor on a cell with the Another aspect of the present invention relates to a method of

WO 01/71022 - 36 -

PCT/US01/08729

compound in a manner effective to enhance LPA receptor-induced proliferation of the cell.

In addition to the roles that LPA plays in modulating cancer cell activity, there is strong evidence to suggest that LPA also has a physiological role in natural wound healing. At wound sites, LPA derived from activated platelets is believed to be responsible, at least in part, for stimulating cell proliferation at the site of injury and inflammation possibly in synchronization with other platelet-derived factors (Balazz et al., 2000). Moreover, LPA by itself stimulates platelet aggregation, which may in turn be the factor that initiates an element of positive feedback to the initial aggregatory response (Schumacher et al., 1979; Tokumura et al., 1981; Gerrard et al., 1979; Simon et al., 1982).

Due to the role of LPA in cell proliferation, compounds having LPA receptor agonist activity can be used in a manner effective to promote wound healing. Accordingly, another aspect of the present invention relates to a method of treating a wound. This method is carried out by providing a compound of the present invention which has activity as an agonist of an LPA receptor and delivering an effective amount of the compound to a wound site, where the compound binds to LPA receptors on cells that promote healing of the wound, thereby stimulating LPA receptor agonist-induced cell proliferation to promote wound healing.

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The primary goal in the treatment of wounds is to achieve wound closure. Open cutaneous wounds represent one major category of wounds and include burn wounds, neuropathic ulcers, pressure sores, venous stasis ulcers, and diabetic ulcers. Open cutaneous wounds routinely heal by a process which comprises six major components: i) inflammation, ii) fibroblast proliferation, iii) blood vessel proliferation, iv) connective tissue synthesis v) epithelialization, and vi) wound contraction. Wound healing is impaired when these components, either individually or as a whole, do not function properly. Numerous factors can affect wound healing, including malnutrition, infection, pharmacological agents (e.g., actinomycin and steroids), diabetes, and advanced age (see Hunt and Goodson, 1988).

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Phospholipids have been demonstrated to be important regulators of cell activity, including mitogenesis (Xu et al., 1995b), apoptosis, cell adhesion, and regulation of gene expression. Specifically, for example, LPA elicits growth factor-like effects on cell proliferation (Moolenaar, 1996) and cell migration (Imamura et al. 1993). It has also been suggested that LPA plays a role in wound healing and regeneration (Tigyi and Miledi, 1992).

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In general, agents which promote a more rapid influx of fibroblasts, endothelial and epithelial cells into wounds should increase the rate at which wounds

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WO 01/71022 -37 -

heal. Compounds of the present invention that are useful in treating wound healing can be identified and tested in a number of in vitro and in vivo models.

It vitro systems model different components of the wound healing process, for example the return of cells to a "wounded" confluent monolayer of tissue culture cells, such as fibroblasts (Verrier et al., 1986), endothelial cells (Miyata et al., 1990) or epithelial cells (Kartha et al., 1992). Other systems permit the measurement of endothelial cell migration and/or proliferation (Muller et al., 1987; Sato et al., 1988).

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In vivo models for wound healing are also well-known in the art including wounded pig epidermis (Ohkawara et al., 1977) or drug-induced oral mucosal lesions in the hamster cheek pouch (Cherrick et al., 1974).

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The compounds of the present invention which are effective in wound healing can also be administered in combination, i.e., in the pharmaceutical composition of the present invention or simultaneously administered via different routes, with a medicament selected from the group consisting of an antibacterial agent an antiviral agent, an antifungal agent, an antiparasitic agent, an antiinflammatory agent, an analgesic agent, an antipruritic agent, or a combination thereof.

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For wound healing, a preferred mode of administration is by the topical route. However, alternatively, or concurrently, the agent may be administrated by parenteral, subcutaneous, intravenous, intramuscular, intraperitoneal or transfermal routes. Alternatively, or concurrently, administration may be by the oral route. The dosage administrated will be dependent upon the age, health, and weight of the recipient, kind of concurrent treatment, if any, frequency of treatment, and the nature of the effect desired.

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For the preferred topical applications, especially for treatment of humans and animals having a wound, it is preferred to administer an effective amount of a compound according to the present invention to the wounded area, e.g., skin surfaces. This amount will generally range from about 0.001 mg to about 1 g per application, depending upon the area to be treated, the severity of the symptoms, and the nature of the topical vehicle employed. A preferred topical preparation is an ointment wherein about 0.01 to about 50 mg of active ingredient is used per ml of ointment base, such as PEG-1000.

PCT/US01/08729

-38-

EXAMPLES

The following examples are intended to illustrate, but by no means are intended to limit, the scope of the present invention as set forth in the appended

Materials and Methods

A Thomas-Hoover capillary melting point (mp) apparatus was used to measure all melting points (mps), which were uncorrected.

¹H and ¹³C nuclear magnetic resonance (NMR) spectra were recorded expressed as parts per million (ppm) relative to tetramethylsilane (TMS). Peaks are on a Bruker AX 300 spectrometer (300, 75.5 MHz). Chemical shift values (8) are abbreviated as follows: s - singlet; d - doublet; t - triplet; q - quartet; bs - broad singlet; m - multiplet.

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Spectra for phosphorous-31 are reported as parts per million () relative to 0.0485 M carbon-13 are reported as parts per million () relative to tetramethylsilane (TMS). Proton, carbon-13, and phosphorous-31 magnetic resonance spectra were obtained on a Bruker AX 300 spectrometer. Chemical shifts for proton and riphenylphosphate in acctone-ds at = 0 ppm.

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infrared (IR) spectra were recorded on Perkin Elmer System 200-FTIR.

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Mass spectra (MS) were recorded on either a Bruker Esquire AG or a Interface (BSI) either in the positive or negative mode. Spectral data were consistent Bruker Bsquire LC/MS spectrometer by direct infusion utilizing the Blectrospray with assigned structures.

Elemental analysis was performed by Atlantic Microlabs, Inc. (Norcross, GA), and values found are within ±0.4% of the theoretical values.

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Silica gel (Merck, 230-400mesh or 200-425 mesh, 60A°) was used for Analytical TLC was performed on Sigma-Aldrich silica gel 60 F 254 flash column chromatography.

TLC sheets with aluminum backings (thickness 200 or 250 microns).

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was distilled from calcium hydride (CaH2). All the mono glycerides were from Nu-All reagents, solvents, and chromatography media, unless otherwise Fisher Scientific (Pittsburgh, PA), or Sigma Chemical Co. (St. Louis, MO) without metal with benzophenone as an indicator. Anhydrous methylene chloride (CH₂Cl₂) further purification. Tetrahydrofuran (THF) was dried by distillation from sodium noted, were purchased from either Aldrich Chemical Company (Milwaukee, WI), Check -Prep (Minneapolis, MN). 6-Boc-L-serine was purchased from Fluka.

32

WO 01/71022

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PCT/US01/08729

Fatty acid-free bovine serum albumin (BSA). Prior to use, LPA was complexed, at a solution containing 1 mM EGTA. Aliquots of all the other lipids were dissolved in All lipids were purchased from Avanti Polar Lipids (Alabaster, AL). 1:1 ratio molar ratio, with 1 mM BSA dissolved in Ca2* -free Hanks' balanced salt MeOH and mixed with LPA prior to application, or as otherwise indicated.

Cytofectene transfection reagent was from Bio-Rad (Hercules, CA). Fura-2 AM was from Molecular Probes (Eugene, OR). Culture media, fetal bovine serum (FBS), and G418 were obtained from Cellgro (Herndon, VA).

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RH7777 cells, stably expressing human Edg-4, were kindly provided by Invitrogen, Carlsbad, CA), were a generous gift from Dr. Junken Aoki (University of Dr. Kevin Lynch (University of Virginia, Charlottesville, VA). Flag-tagged cDNA's Tokyo, Tokyo, Japan). RH7777 and NIH3T3 cells were obtained from the American Dulbecco's Modified Eagle's Medium (DMEM) containing 10% FBS and 2 mM glutamine. Oocytes were obtained from adult Xenopus laevis frogs as previously Type Culture Collection (Manassas, VA). HEY cells were provided by Dr. Lisa emings (University of Tennessee, Memphis). All cell lines were maintained in encoding human Edg-4 and -7 inserted into the pCDNA3 expression plasmid described (Tigyi et al., 1999).

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Stable transfection

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protocol. Transfected cells were selected in DMEM containing 10% FBS and 1 mg/ml numan Edg-2, Edg-4, or Edg-7 and then were subcloned into the pCDNA3 expression RH7777 cells were transfected with the cDNA constructs encoding geneticin. Resistant cells were collected and subcloned by limiting dilution. The esulting clones were then screened using functional assays and RT-PCR analysis. vector using the Cytofectene transfection reagent according to the manufacturers' Data are representative of three individual clones.

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Transient transfection

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containing 10% FBS. The next day, the cells were rinsed with DMEM and serum was Sysofectene. The cells were then rinsed twice with DMEM and cultured in DMEM (Bellco, Vineland, NJ) one day prior to transfection. The following day, cells were ransfected overnight (16-18 hr) with 1 µg of plasmid DNA mixed with 6 µl of RH7777 cells were plated on polylysine-coated glass coverslips withdrawn for a minimum of $2 \, \mathrm{lrr}$ prior to monitoring intracellular Ca $^{2+}$

76 01/71022 - 40 -

PCT/US01/08729

Measurement of intracellular Ca2+ and data analysis

Changes in intracellular Ca²⁺ were monitored using the fluorescent Ca²⁺ indicator Pura-2 AM as previously described (Tigyi et al., 1999). Dam points from the intracellular Ca²⁺ measurements represent the total peak area of the Ca²⁺ transients elicited, as determined by the FLWinLab software (Perkin-Elmer, Wellesley, MA). Data points represent the average of at least 3 measurements ± standard deviation. The significance of the data points was determined using the students t-test and values were considered significant at p < 0.05.

Electrophysiological recording in Xenopus occytes

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Oscillatory Cl' currents, elicited by LPA, were recorded using a twoelectrode voltage clamp system as previously described (Tigyi et al., 1999).

RT-PCR analysis of Edg and PSP24 mRNA

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The identification of Edg and PSP24 receptor mRNA by RT-PCR was performed as previously described (Tigyi et al., 1999), using the following oligonucleotide sequences:

EDQ-1

forward primer 5'-s1TCATCGTCCGGCATTACAACTA-3' (SEQ. ID No. 9);
reverse primer 5'-QAGTGAGCTTGTAGGTGGTG₃₅₁-3' (SEQ. ID No. 10);
EDG.2

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forward primer 5'-syAGATCTGACCAGCCGACTCAC-3' (SEQ. ID No. 11); reverse primer 5'-GTTGGCCATCAAGTAATAAATA₄₂₇-3' (SEQ. ID No. 12); EDG-3

forward primer 5'-137CTTGGTCATCTGCAGCTTCATC-3' (SEQ. ID No. 13); reverse primer 5'-TGCTGATGCAGAAGGCAATGTA397-3' (SEQ. ID No. 14); EDG-4

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forward primer 5'-sa/CTGCTCAGCCGCTCCTATTTG-3' (SEQ. ID No. 15);
reverse primer 5'-AGGAGCACCCACAAGTCATCAG1185-3' (SEQ. ID No. 16);
EDG-5

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forward primer 5'-1;ATGGGCAGCTTGTACTCGGAG-3' (SEQ. ID No. 17);
reverse primer 5'-CAGCCAGCAGACGATAAAGAC_{720'}-3' (SEQ. ID No. 18);
proc.4

forward primer 5'-2s0TGAACATCACGCTGAGTGACCT-3' (SEQ. ID No. 19) reverse primer 5'-GATCATCAGCACCGTCTTCAGC₇₈₀-3' (SEQ. ID No. 20); EDG-7

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forward primer 5'-91AGCAACACTGATACTGTCGATG-3' (SEQ, ID No. 21); reverse primer 5'-GCATCCTCATGATTGACATGTG46-3' (SEQ, ID No. 22);

WO 01/71022

-41-

PCT/US01/08729

EDG-8

forward primer 5'-seATCTGTGCGCTCTATGCAAGGA-3' (SEQ. ID No. 23); reverse primer 5'-GGTGTAGATGATAGGATTCAGCA₁₁₆₁ -3' (SEQ. ID No. 24); PSP24

forward primer 5'-220CTGCATCATCGTGTACCAGAG-3' (SEQ. ID No. 25); and reverse primer 5'-ACGAACTCTATGCAGGCCTCGC1184-3' (SEQ. ID No. 26).

Cell proliferation assay Proliferation of NIH3T3 cells was assessed by direct cell counting as

previously described (Tigyi et al., 1999). NIH3T3 cells were plated in 24-well plates at a density of 10,000 cells/well, in DMEM containing 10% FBS. The following day, the cells were rinsed and serum starved in DMEM for 6 hr. Lipids were then added for 24 hr. Cell numbers were determined by counting in a Coulter counter (Coulter Electronics, Hialeah, FL).

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Incorporation of 3H-thymidine

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The incorporation of ³H-thymidine into RH7777 cells was determined as previously described (Tigyi et al., 1994).

Example 1. Synthesis of N-(tert-butoxycarbonyl)-L-serine β-lactone, Intermediate Compound 25

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of 30 min. After the addition was complete, the mixture was stirred until a milky oil at 30 °C.: The oil was then treated with 25% EtOAc/hexanes (100 ml), the resulting mixture was stirred for 2 hrs and concentrated on the rotary evaporator to pale yellow mixture. The mixture was stirred overnight at -78 °C under argon and allowed to white paste was obtained (ca. 30-40 min). A solution of N-(tert-butoxycarbonyl)-Lazodicarboxylate (DEAD) (6.2 ml, 39.9 mmol) was added with a syringe over a period triphenylphosphine (Ph₃P) (10 g, 38 mmol, dried over P₂O₅ under vacuum for 72 hrs) white solid was removed by filteration and washed with 25% EtOAc/hexanes (2×70 °C). After 30 min (ca) the ice bath was replaced with a water bath, and the reaction warm to 0 °C (the flask was placed in an ice bath when the temperature reached -10 distilled THF (75 ml) was added dropwise over a period of 45 min to the reaction serine (24) (7.79 g, 38 mmol, dried over P₂O₅ under vacuum for 72 hrs) in freshly ice-acetone bath) under argon. With vigorous stirring, freshly distilled diethyl and freshly distilled THF (190 ml). The solution was cooled and stirred at ~78 °C (dry cooled to room temperature under Argon (Ar) before use. To the flask were added thermometer and a 100 ml dropping funnel. All glassware were flame-dried and A 500 ml three-neck flask was equipped with a low temperature

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- 42 -

ml), the combined filtrate was concentrated, and the residual oil subjected to flash chromatography on silica gel with 25% (500 ml) and 30% (1500 ml) EtOAc/hexanes, successively. Appropriate fractions were combined to afford 3.4 g (47%) of 25 as a white solid: mp 119-121 °C dec (Lit. 119.5-120.5 °C dec); ¹H NMR (CD₂Cl₂) § 1.44 (8, 9H), 4.38-4.42 (m, ZH), 4.96-5.03 (q, J₁= 6.1 Hz, J₂=12.5 Hz, 1H), 5.39 (s, br, 1H); ¹³C NMR (CD₂Cl₂) d 28.31, 60.01, 66.63, 81.50, 155.01, 169.94; IR (KBr) 3361, 2978, 1843, 1680, 1533, 1370, 1292 cm⁻¹; Anal. Caled. for C₂H₁₃NO₄: C, 51.33; H, 6.94; N, 7.50. Found: C, 51.41; H, 7.01; N, 7.51.

Example 2 - Synthesis of Compounds 26-34

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The glassware used were flame-dried and cooled to room temperature under argon atmosphere. The reaction was carried out in argon atmosphere. THF was freshly distilled prior to use.

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Compound 26: tert-Butyl N-[1-(hydroxymethyl)-2-(nonylamino)-2-oxoethyl]carbamate

To a solution of decyl amine (490 mg, 3.20 mmol) in THF (60 ml), N- (terr-butoxycarbonyl)-L-serine β-lactone (300 mg, 1.60 mmol) was added, and the mixture was refluxed overnight under argon. The reaction mixture was concentrated on a rotary evaporator. The residue was subjected to flash column chromatography, eluting with EtOAc/hexanes of various compositions.

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Appropriate fractions were pooled, and concentrated to dryness in vecuo to afford 290 mg (52%) of 26 as a white waxy powder: mp 50-52°C; ¹H NIMR (CDCI₃) 8 0.88 (t, 1=6.4 Hz, 3H), 1.26 (s, 14H), 1.46 (a, 9H), 3.04 (bs, 1H), 3.16-3.34 (m, 2H), 3.63 (m, 1H), 4.06-4.15 (m, 2H), 5.37 (bs, 1H), 3.04 (bs, 1H); ¹³C NIMR (CDCI₃) 8 1409, 22.65, 26.80, 28.27, 29.24, 29.27, 29.37, 29.50, 29.51, 31.86, 39.43, 54.34, 62.87, 77.20, 80.34, 171.52; IR (KBt) 32.82, 3098, 2929, 2856, 1666, 1547, 1467, 1369, 1300, 1248, 1179 cm⁻¹; Anal. Calcd. for C₁₆H₂₂N₂O₄: C, 62.76; H, 10.53; N, 8.13. Found: C, 63.00; H, 10.46; N, 7.98.

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Compound 27: terr-Butyl N-{1-{hydroxymethyl}-2-oxo-2-(tetradecylamino)ethyl]carbamate

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To a solution of tetradecyl amine (273 mg, 1.28 mmol) in THF (40 ml), N-(terr-butoxycarbonyl)-L-serine β-lactone (200 mg, 1.06 mmol) was added, and the mixture was refluxed overnight under argon. The reaction mixture was concentrated on a rotary evaporator. The residue was subjected to flash column chromatography, eluting with EtOAchexanes of various compositions.

33

WO 01/71022

- 43 - PCT/US01/08729

Appropriate fractions were pooled, and concentrated to dryness in vacuo to afford 245 mg (57%) of 27 as a white powder: mp 59-62°C; ¹H NMR (CDCl₃) 8 0.88 (t, 1=6.3 Hz, 3H), 1.25 (s, 24H), 1.45 (s, 9H), 3.15-3.36 (m, 3H), 3.65 (m, 1H), 4.07-4.13 (m, 2H), 5.60-5.63 (m, 1H), 6.72 (bs, 1H); ¹³C NMR (CDCl₃) 8 14.10, 22.66, 26.81, 27.99, 28.27, 29.25, 29.33, 29.37, 29.50, 29.57, 29.65, 31.90, 39.47, 54.58, 62.87, 77.20, 80.52, 156.34, 171.37; R (KBr) 3345, 2920, 2852, 1708, 1688, 1655, 1637, 1572, 1529, 1472, 1248, 1173 cm⁻¹; Anal. Calcd. for C₂H₄M₂O₄. C, 65.96; H, 11.07; N, 6.99. Found: C, 66.04; H, 11.17; N, 6.96.

Compound 28: tert-Butyl N-[1-(hydroxymethyl)-2-(octadecylamino)-2-oxoethyl]carbamate

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To a solution of octadecyl amine (516 mg, 2.08 mmol) in THF (60 ml), N-(rerr-butoxycarbonyl)-L-serine β-lactone (300 mg, 1.60 mmol) was added, and the mixture was refluxed overnight under argon. The reaction mixture was concentrated on a rotary evaporator. The residue was subjected to flash column chromatography, cluting with EtOAchtexanes of various compositions.

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Appropriate fractions were pooled, and concentrated to dryness in vacuo to afford 300 mg (41%) of 28 as white powder: mp 69-71°C; ¹H NMR (CDCl₃) δ 0.88 (t, J=6.3 Hz, 3H), 1.25 (s, 30H), 1.46 (s, 9H), 3.03 (bs, 1H), 3.16-3.34 (m, 2H), 3.63 (m, 1H), 4.05-4.21 (m, 2H), 5.64 (bs, 1H), 6.62 (bs, 1H); ¹³C NMR (CDCl₃) δ 14.10, 22.68, 26.81, 28.28, 29.25, 29.51, 29.59, 29.59, 29.69, 31.91, 39.43, 54.29, 62.87, 77.20, 171.53; R (KBr) 3345, 2919, 2852, 1687, 1636, 1570, 1528, 1473, 1305, 1173 cm⁻¹; Anal. Calcd. for C₂₆H₂NλO₄ · 0.2C₄H₄O₅: C, 67.86; H, 11.39; N, 5.91. Found: C, 67.59; H, 11.46; N, 6.1.

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Compound 29: tert-Butyl N- [1-(hydroxymethyl)-2-oxo-2-[4-(tetredecyloxy)anilino]ethyl) carbamate

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To a solution of 4-(tetradecyloxy)aniline (150 mg, 0.490 mmol) in THF (40 ml), N-(terr-butoxycarbony)-L-serine β-lactone (91 mg, 0.490 mmol) was added, and the mixture was refluxed for 48 hrs under argon. The reaction mixture was concentrated on a rotary evaporator. The residue was subjected to flash column chromatography (twice), eluting with EKOAc/hexanes of various compositions.

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Appropriate fractions were pooled, and concentrated to dryness in vacuo to afford 110 mg (45%) of 29 as a white powder: mp 92-94 °C; ¹H NMR (CDCl₃) 8 0.87 (t, J=6.6 Hz, 3H), 1.25 (s, 22H), 1.48 (s, 9H), 1.76 (m, 2H), 3.67-3.72 (dd, J₁=4.9 Hz, J₂=7.2 Hz, 1H), 3.92 (t, J=6.5 Hz, 2H), 4.23-4.26 (m, 2H), 5.65 (bs, 1H), 6.83-6.87 (m, J₆=8.9 Hz, 2H), 7.36-7.40 (m, J₆=8.9 Hz), 8.6 (bs, 1H); ¹³C NMR (CDCl₃) 8 14.10, 22.69, 26.01, 28.28, 29.25, 29.34, 29.39, 29.56, 29.58, 29.64, 31.91,

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PCT/US01/08729

0.05CHCl₃: C, 67.56; H, 9.71; N, 5.62. Found: C, 67.80; H, 9.67; N, 5.60. 2920, 2852, 1658, 1514, 1472, 1238, 1174 cm⁻¹; Anal. Calcd. for C₂₅H₄₅N₂O₅. 62.53, 68.30, 77.20, 111.17, 114.81, 121.70, 130.25, 156.22, 169.78; IR (KBr) 3304,

Compound 30: tert-Butyl N-[1-(hydroxymethyl)-2-(4methoxyanilino)-2-oxoethyl]carbamate

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on a rotary evaporator. The residue was subjected to flash column chromatography, cluting with EtOAc/hexanes of various compositions. mixture was refluxed overnight under argon. The reaction mixture was concentrated (tert-butoxycarbonyl)-L-serine β -lactone (151 mg, 0.8 mmol), was added, and the To a solution of p-anisidine (100 mg, 0.8 mmol) in THF (20 ml) N-

Anal. Calcd. for C15H2N2O5: C, 58.05; H, 7.15; N, 9.03. Found: C, 58.04; H, 7.17; N ¹³C NMR (CDCl₃) 8 28.29, 54.96, 55.47, 62.54, 81.00, 114.18, 121.78, 130.45, 5.68 (bs, 1H); 6.83-6.88 (m, J₀=9 Hz, 2H), 7.37-7.42 (m, J₀=9 Hz, 2H), 8.61 (bs, 1H); NMR (CDCl₃), 8 1.48 (s, 9H), 3.68-3.73 (m, 1H), 3.80 (s, 3H), 4.24-4.27 (m, 2H), 156.64, 156.98, 169.59; IR (KBr) 3340, 2978, 1673, 1603, 1516, 1298, 1238, cm⁻¹; CHClyhexanes to afford 135 mg (54%) of 30 as a white powder: mp 109-111°C; 'H Appropriate fractions were pooled, and were crystallized from

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Compound 31: tert-Butyl N-{1-(hydroxymethyl)-2-0x0-2-[3-(tetredecyloxy)anilino]ethyl} carbamate

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chromatography, cluting with EtOAc/hexanes of various compositions. and the mixture was refluxed for 48 hrs under argon. The reaction mixture was (25 ml), N-(tert-butoxycarbonyl)-L-scrine β-lactone (91 mg, 0.490 mmol) was added, concentrated on a rotary evaporator. The residue was subjected to flash column To a solution of 3-(tetradecyloxy)amiline (179 mg, 0.588 mmol) in THF

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(CDCl₃) 8 0.88 (t, J=6.6 Hz, 3H), 1.26 (s, 22H), 1.48 (s, 9H), 1.76 (m, 2H), 3.67-3.73 vacuo to afford 105 mg (43%) of 31 as a white powder: mp 70-72 °C; ¹H NMR Calcd. for C28H49N2O3 · 0.05CHCl3: C, 67.56; H, 9.71; N, 5.62. Found: C, 67.44; H. 138.54, 159.75; IR (KBr) 3368, 2918, 2851, 1679, 1618, 1498, 1472, 1286 cm 1; Anai 29.39, 29.60, 29.66, 31.92, 62.38, 68.07, 77.20, 106.22, 111.10, 111.92, 129.67, (dd, J₁=5.1 Hz, J₂= 6.9 Hz, 1H), 3.93 (t, J=6.5 Hz, 2H), 4.23-4.26 (m, 2H), 5.66 (bs, 1H), 8.75 (bs, 1H); ¹³C NMR (CDCl₃) 8 14.11, 22.68, 26.02, 28.28, 29.23, 29.35, 1H), 6.64-6.68 (m, 1H), 6.93-6.96 (m, 1H), 7.19 (t, J_o=8.1 Hz, 1H), 7.23 (t, J_o=2 Hz, Appropriate fractions were pooled, and concentrated to dryness in

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WO 01/71022

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PCT/USU1/08729

Compound 32: tert-Butyl N-[1-(hydroxymethyl)-2-(3methoxyanilino)-2-oxoethyljcarbamate

on a rotary evaporator. The residue was subjected to flash column chromatography mixture was refluxed overnight under argon. The reaction mixture was concentrated eluting with EtOAc/hexanes of various compositions. (tert-butoxycarbonyl)-L-scrine β -lactone (200 mg, 1.06 mmol) was added, and the To a solution of m-anisidine (171 mg, 1.38 mmol) in THF (30 ml), N-

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6.96-6.99 (m, 1H), 7.21 (m, I₀=8.1 Hz, 1H), 7.24 (m, 1H), 8.79 (bs, 1H); ¹³C NMR yellow oil; 1 H NMR (CDCl₃), δ 1.48 (s, 9H), 3.68-3.73 (dd, J_1 =4.8 Hz, J_2 =6.9 Hz, 1H), 3.75 (s, 3H), 4.22-4.25 (d, J=10.23 Hz, 2H), 5.66 (bs, 1H), 6.66-6.69 (m, 1H), 138.63, 160.19, 169.89. (CDCl₃) 6 28.28, 29.68, 55.30, 62.39, 77.20, 81.11, 105.67, 110.55, 112.15, 129.73 Appropriate fractions were pooled, to afford 154 mg (46%) of 32 as a

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Compound 33: tert-Butyl N-{1-(hydroxymethyl)-2-oxo-2-[2-(tetredecyloxy)anilinojethyl) carbamate

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chromatography, eluting with EtOAc/hexanes of various compositions. concentrated on a rotary evaporator. The residue was subjected to flash column and the mixture was refluxed for 48 hrs under argon. The reaction mixture was (25 ml), N-(tert-butoxycarbonyl)-L-serine β-lactone (102 mg, 0.545 mmol) was added To a solution of 2-(tetradecyloxy)aniline (200 mg, 0.654 mmol) in THF

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31.91, 55.73, 63.03, 68.71, 77.20, 111.06, 119.86, 119.86, 120.78, 124.21, 127.27, 6.9 Hz, 1H), 3.93 (t, J=6.5 Hz, 2H), 4.23-4.26 (m, 2H), 5.66 (bs, 1H), 6.64-6.68 (m, J=6.6~Hz, 3H), 1.26 (s, 22H), 1.48 (s, 9H), 1.76 (m, 2H), 3.67-3.73 (dd, $J_1=5.1~Hz$, $J_2=$ 147.75, 157.22, 169.25. 13C NMR (CDCl₃) & 14.10, 22.68, 25.88, 28.30, 29.17, 29.35, 29.58, 29.64, 29.68, 1H), 6.93-6.96 (m, 1H), 7.19 (t, J₆=8.1 Hz, 1H), 7.23 (t, J_m=2 Hz, 1H), 8.75 (bs, 1H); vacuo to afford 33 mg (< 10%) of 33 as a yellow oil: ${}^1\!H$ NMR (CDCl₃) δ 0.88 (t, Appropriate fractions were pooled, and concentrated to dryness in

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Compound 34: tert-Butyl N-[1-(hydroxymethyl)-2-(2methoxyanilino)-2-oxosthyl]carbamate

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on a rotary evaporator. The residue was subjected to flash column chromatography, mixture was refluxed for 48 hrs under argon. The reaction mixture was concentrated eluting with EtOAc/hexanes of various compositions (tert-butoxycarbonyl)-L-serine β -lactone (200 mg, 1.06 mmol) was added, and the To a solution of o-anisidine (238 mg, 1.93 mmol) in THF (30 ml), N-

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CHClyhexanes to afford 150 mg (45%) of 34 as a yellow powder: mp 92-94°C; 'H Appropriate fractions were pooled, and crystallized from

- 46 -

NMR (CDCl₃), δ 1.49 (s, 9H), 3.87 (s, 3H), 3.73-3.83 (m, 1H), 4.21,-4.34 (m, 2H), 5.64 (bs, 1H), 6.86-6.97 (m, 2H), 7.03-7.09 (m, J₀=7.80 Hz, J_m=1.8 Hz, 1H), 8.28-8.31 (dd, J₀=8.9 Hz, J_m=1.5 Hz, 1H) 8.9 (bs, 1H); ¹³C NMR (CDCl₃) δ 28.28, 55.73, 62.87, 80.65, 110.14, 120.03, 120.97, 124.30, 127.13, 148.33, 169-43; R. (KBr) 33225, 3319, 2982, 1672, 1653, 1548, 1528, 1465, 1256, 1160, 1006 cm⁻¹; Anal. Calcd. for C₁Hz₂N₂O₂: C, 58.05; H, 7.15; N, 9.03. Found: C, 58.04; H, 7.07; N, 8.85.

Example 3 - Synthesis of Compounds 35-43

Compound 35: N-1-nonyl-2-amino-3-hydroxypropanamide trifluoroacetate

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To a cooled (0 °C, ice bath) solution of 26 (20 mg, 0.0580 mmol) in CH₂Cl₂ (1 ml), TFA (1 ml) was added dropwise under argon atmosphere. After the addition was complete, the reaction was allowed to sit at r.t., for 3 hrs, concentrated under reduced pressure at room temperature, and dried on a vacuum pump to give 35 as a white solid 19 mg (95%): mp 168-170°C; ¹H NMR (CD₃OD), 5 0.88 (t, J=6.3 Hz, 3 H), 1.27 (s, 14H), 1.50 (m, 2H), 3.20 (t, J=6.0 Hz, 2H), 3.70-3.78 (m, 1H), 3.81-3.88 (m, 2H); ¹³C NMR (CD₃OD) 5 14.44, 23.74, 27.96, 30.30, 30.42, 30.47, 30.70, 30.73, 30.78, 30.80, 33.10, 40.71, 56.30, 61.77, 167.97; IR (KBr) 3280, 2919, 2850, 1654, 1573, 1464, 1231, 1141, 1089, 1059, cm⁻¹. Anal. Calcd. for C₁₃H₂₈N₂O₂·CF₃COOH: C, 50.27; H, 8.16; N, 7.82. Found: C, 50.15; H, 8.30; N, 7.95.

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Compound 36: N-1-tetradecyl-2-amino-3-hydroxypropanamide trifluoroacetate

To a cooled (0 °C, ice bath) solution of 27 (50 mg, 0.124 mmol) in CH₂Cl₂ (1.5 ml), TFA (1.5 ml) was added dropwise under argon atmosphere. After the addition was complete, the reaction was allowed to stir at r.t. for 3 hrs, concentrated under reduced pressure at room temperature, and dried on a vacuum pump to give 36 as a white solid 48 mg (94%): mp 168-171°C; ¹HNMR (CD₂OD), δ 0.89 (t, 1=6.3 Hz, 3H), 1.28 (a, 22H), 3.22 (t, 1=6.0 Hz, 2H), 3.73-3.80 (m, 1H), 3.84-3.91 (m, 2H), ¹³C NMR (CD₃OD) δ 14.43, 23.73, 27.95, 30.29, 30.41, 30.47, 30.69, 30.73, 30.78, 30.89, 31.08, 40.71, 56.29, 61.77, 167.99; IR (KBr) 3277, 2919, 2850, 1656, 1573, 1464, 1231, 1141, 1089, 1059 cm⁻¹; Anal. Calcd. for C₁₇H₃8N₂O₂-CF₃COOH: C, 55.06, H, 9.00; N, 6.76. Found: C, 54.94; H, 8.99; N,

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WO 01/71022

- 47 -

PCT/US01/08729

Compound 37: N-1-octadecyl-2-amino-3-hydroxypropanamide trifluoroacetate

To a cooled (0 °C, ice bath) solution of 28 (25 mg, 0.0547 mmol) in CH₂Cl₂ (1 ml), TFA (1 ml) was added dropwise under argon atmosphere. After the addition was complete, the reaction was allowed to stir at r.t., for 3 hrs, concentrated under reduced pressure at room temperature, and dried on a vacuum pump to give 37 as a white solid 23 mg (92%): mp 170-172°C, ¹H NMR (CD₃OD) § 0.89 (t, J=6.4 Hz, 3H), 1.27 (s, 30H), 1.49-1.54 (m, 2H), 3.22 (t, J=7.0 Hz, 2H), 3.74-3.81 (m, 1H), 3.83-3.91 (m, 2H); ¹C NMR (CD₃OD) § 14.43, 23.74, 27.95, 30.30, 30.41, 30.47, 30.69, 30.78, 33.07, 40.71, 56.30, 61.77, 167.97; IR (KBr) 3276, 2919, 2850, 1657, 1468, 1207, 1181, 1138, 1059 cm⁻¹; Anal. Calcd. for C₂H4₄M₂O₂-CP₂COOH 0.15CH₂Cl₂: C, 57.53; H, 9.45; N, 5.80. Found: C, 57.45; H, 9.55; N, 5.81.

Compound 38: N-1-[4-(tetradecyloxy)phenyl]-2-amino-3hydroxypropanamide trifinoroacetate

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To a cooled (0 °C, ice bath) solution of 29 (54 mg, 0.110 mmol) in CH₂Cl₂ (0.050 ml), TFA (0.050 ml) was added dropwise under argon atmosphere. After the addition was complete, the reaction was allowed to stir at r.t., for 3 hrs, concentrated under reduced pressure at room temperature, and then, dried on a vacuum pump to give 38 as a white solid 55 mg (99%); mp 135-139 °C; ¹H NNR (CD₂)OD), 8 (0.89 (t, 1=6.3 Hz, 2H), 1.28 (s, 21H), 1.43 (m, 2H), 1.74 (m, 1=6.5 Hz, 2H), 13.66-4.03 (m, 5H), 6.84-6.88 (m, 1₀=9.0 Hz, 2H), 7.41-7.47 (m, 1₀=9.0 Hz, 2H); ¹³C NNR (CD₂)OD) § 14.42, 23.72, 30.41, 30.46, 30.50, 30.67, 30.74, 33.06, 56.81, 61.72, 69.26, 115.71, 122.96, 131.84, 157.80, 166.06; IR (KBr) 3281, 2920, 2852, 1672, 1604, 1559, 1515, 1240, 1210, 1132 cm⁻¹, Anal. Calcd. for C₂₂H₄₀N₂O₃. CF₃COOH: C,

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Compound 39: N-1-(4-methoxyphenyl)-2-amino-3hydroxypropanamide trifluoroacetate

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19.27; H, 8.16; N, 5.53. Found: C, 59.48; H, 8.09; N, 5.49.

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To a cooled (0 °C, ice bath) solution of 30 (50 mg, 0.161 mmol) in CH₂Ch₂ (0.049ml), TFA (0.049 ml) was added dropwise under argon atmosphere. After the addition was complete, the reaction was allowed to stir at r.t., for 3 hrs, concentrated under reduced pressure at r.t., and concentrated to dryness in vacuo to give 39 as a white solid 50 mg (96%); mp 182-183°C dec; ¹H NMR (CD₂OD), 6 3.76 (s, 3H), 3.87-3.94 (m, 1H), 3.97-4.04 (m, 2H), 6.85-6.91 (m, 1₆-9.1 Hz, 2H), 7.44-7.49 (m, 1₆-9.0 Hz, 2H); ¹³C NMR (CD₅OD) 5 55.86, 56.80, 61.73, 115.07, 122.95, 131.99, 158.31, 166.10; IR (KBr) 32.78, 3099, 2964, 1673, 1562, 1517, 1196, 1131, cm⁻¹; Anal Calcd. for C₁₀H₁Mh₂O₂ · CF₂COOH: C, 44.45; H, 4.66; N, 8.64. Found: C, 44.31; H, 4.67; N, 8.58.

- 48 -

WO 01/71022

Compound 40: N-1-[3-(tetradecyloxy)phenyl]-2-amino-3hydroxypropanamide trifluoroacetate

To a cooled (0 °C, ice bath) solution of 31 (45 mg, 0.091 mmol) in CH₂Ch (0.062 ml), TFA (0.062 ml) was added dropwise under argon atmosphere. After the addition was complete, the reaction was allowed to stir at r.t. for 3 hrs, concentrated under reduced pressure at room temperature, and dried on a vacuum pump to give 40 as a yellowish green solid 45 mg (99%): mp 115-119 °C; ¹H NMR (CD₃OD), 8 0.89 (s, J=6.5Hz, 3H), 1.28 (s, 21H), 1.43 (m, 2H), 1.75 (m, J=6.5 Hz, 2H), 3.8-3.93 (m, 4H), 4.01-4.05 (m, 1H), 6.67-6.71 (m, 1H), 7.04-7.07 (m, 1H), 7.20 (t, J₀=8.1 Hz, 1H), 7.28 (t, J_m=2.1 Hz, 1H), 11°C NMR (CD₃OD) 814-44, 23.75, 27.18, 30.38, 30.49, 30.52, 30.73, 30.78, 33.09, 56.96, 61.66, 69.05, 107.71, 111.75, 113.16, 130.72, 140.16, 161.07, 166.36; IR (KBr) 3266, 2920, 2852, 1676, 1608, 1566, 1496, 1438, 1211, 1130, 1045 cm⁻¹; Anal. Calcd. for C₂₂H₄₀N₂O₃ · CF₃COOH: C, 59.27; H, 8.16; N, 5.53. Found: C, 59.49; H, 8.13; N, 5.41.

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Compound 41: N-1-(3-methoxyphenyl)-2-amino-3hydroxypropanamide triffuoroacetate

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To a cooled (0 °C, ice bath) solution of 32 (120 mg, 0.386 mmol) in CH₂Cl₂ (1 ml), TFA (1 ml) was added dropwise under argon atmosphere. After the addition was complete, the reaction was allowed to stir at r.t., for 3 hrs, concentrated under reduced pressure at r.t., and dried on a vacuum pump to give 41 as a off-white solid 123 mg (98%): mp 137-140 °C; ¹HNMR (CD₃OD), 6 3.77 (s, 3H), 3.88-3.99 (m, 2H), 4.01-4.06 (m, 1H), 6.68-6.71 (m, 1H), 7.02-7.10 (m, 1H), 7.22 (t, I₃=8.1 Hz, 1H), 7.29 (t, I₃=2.1 Hz, 1H); ¹³C NMR (CD₃OD) 8 55.70, 56.94; 61.67, 107.14, 111.11, 113.28, 130.73, 140.22, 161.61, 166.43; IR (KB₁) 3265, 1675, 1609, 1566, 1496, 1433, 1268, 1196, 1044, cm⁻¹; Anal. Caled. for C₁₀H₄M₂O₃ · CF₃COOH: C, 44.45; H, 4.66; N, 8.64. Found: C, 44.52; H, 4.59; N, 8.66.

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Compound 42: N-1-[2-(tetradecyloxy)phenyl]-2-amino-3hydroxypropanamide trifluoroacetate

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To a cooled (0 °C, ice bath) solution of 33 (21 mg, 0.044 mmol) in CH₂Cl₂ (1 ml), TFA (1 ml) was added dropwise under argon atmosphere. After the addition was complete, the reaction was allowed to stir at r.t., for 3 hrs, concentrated under reduced pressure at room temperature, and dried on a vacuum pump to give 42 as a offwhite solid 21 mg (95%): mp 63-66 °C; ¹H NMR (CD₂OD), 8 0.88 (t, 1=6.5 Hz, 3H), 1.27 (s, 21H), 1.46 (m, 2H), 1.83 (m, 1=7.8 Hz, 2H), 3.90-4.07 (m, 4H), 4.18 (t, 1=5.8 Hz, 1H), 6.87-6.93 (m, 1H), 6.99-7.02 (m, 1H), 7.08-7.14 (m, 1H), 7.96-7.99 (m, 1H); 110 NMR (CD₂OD) 8 14.43, 23.73, 27.07, 30.27, 30.48, 30.57, 30.79, 33.07, 56.198,

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WO 01/71022 - 49 -

61.67, 69.84, 112.93, 121.40, 123.38, 126.80, 127.53, 150.93, 166.74; IR (KBr) 3282, 2925, 2851, 1679, 1556, 1496, 1458, 1213, 750, cm⁻¹; Anal. Calcd. for C₂₃H₄₀N₂O₃-CF₃COOH 0.5H₂O: C, 58.24; H, 8.21; N, 5.43. Found: C, 58.59; H, 8.09; N, 5.24

Compound 43: N-1-(2-methoxyphenyl)-2-amino-3hydroxypropanamide trifiuoroacetate

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To a cooled (0 °C, ice bath) solution of 34 (80 mg, 0.257 mmol) in CH₂Cl₂ (1 ml), TFA (1 ml) was added dropwise under argon atmosphere. After the addition was complete, the reaction was allowed to stir at r.t., for 3 hrs, concentrated under reduced pressure at room temperature, and dried on a vacuum pump to give 43 as a off white solid 81 mg (97%): mp 131-133 °C; ¹H NMR (CD₂OD), 6 3.88 (s, 3H), 3.91-4.02 (m, 2H), 4.18-4.22 (m, 1H), 6.89-6.94 (m, 1H), 7.01-7.04 (m, 1H), 7.10-7.16 (t, J₂=8.1 Hz, 1H), 8.00-8.03 (t, J_m=2.1 Hz, 1H); ¹¹C NMR (CD₂OD) 8 56.27, 56.34, 56.47, 61.81, 111.94, 121.52, 123.21, 126.71, 127.54, 151.43, 166.80; IR (KBr) 3271, 1675, 1546, 1499, 1465, 1439, 1268, 1207, 1130, cm¹¹; Anal. Calcd. for C₁₀H₁₄N₂O₃ · CF₃COOH: C, 44.45; H, 4.66; N, 8.64. Found: C, 44.18; H, 4.57; N, 8.59.

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Example 4 - Synthesis of Intermediate Compounds 50-54

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The glassware used is flame-dried and cooled to room temperature under an argon atmosphere. The starting alcohol was washed with anhydrous pyridine (3 times), and dried (high vacuum for 48 hrs). The reaction was carried out in an argon atmosphere. THF and CH₂Cl₂ were freshly distilled prior to their use.

Compound 50: tert-Butyl N-[1-{([di(benzyloxy)phosphoryl] oxy)methyl}-2-(nonylamino)-2-oxoethyl] carbamate

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To the pyridine-washed starting 28 (252 mg, 0.551 mmol) was added 1H-tetrazole (231 mg, 3.31 mmol). To this mixture was added a 1:1 mixture of freshly distilled THF/CH₂Cl₂ (50 ml). After 10 mins, dibenzyldiisopropyl phosphoramidate (1.14 gm, 3.31 mmol) was added, and the reaction was stirred under an argon atmosphere for 90 mins. The TLC of the reaction mixture showed the formation of the product This mixture was cooled to 0 °C (ice bath), and a large excess of peracetic acid was added. The mixture was stirred for another 35 mins, followed by the addition of Na-metablisulfite to quench the excess peracetic acid. The THF and CH₂Cl₂ were removed under reduced pressure. The concentrate was treated with BiOAc (70 ml), and was washed with Na-metablisulfite (2×25 ml), NaHCO₃ (2×30 ml), water (2×30

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concentrated under reduced pressure. The residue was subjected to flash column ml), and brine (2x30 ml). The organic portion was dried over NaSO4, and chromatography, eluting with EtOAc/hexanes of various compositions.

4.09 (m, 1H), 4.31-4.43 (m, 2H), 4.96-5.09 (m, 4H), 5.55 (bs, 1H), 6.33 (bs, 1H) 7.31-128.72, 135.02, 168.50; MS m/z 603 (M-H); IR (KBr) 3349, 2919, 2852, 1717, 1685, 7.39 (m, 10H); ¹³C (CDCl₃) 8 14.09, 22.66, 26.79, 28.25, 29.24, 29.27, 29.42, 29.50, J=6.4 Hz, 3H), 1.25 (bm, 29H), 1.34 (m, 2H), 1.44 (s, 9H), 3.17-3.23 (m, 2H), 4.01vacuo to afford 195 mg (49 %) of 50 as a colorless oil: ¹H NMR (CDCl₃) 8 0.87 (t, Appropriate fractions were pooled, and concentrated to dryness in 29.53, 31.86, 39.68, 66.98, 69.66, 69.73, 77.20, 128.06, 128.10, 128.64, 128.70, . 1654, 1516, 1470, 1457, 1242, 1163, 1037, 1025, 999 cm⁻¹.

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Compound 51: terr-Butyl N-[1-{([di(henzyloxy)phosphoryl] oxy)methyl}-2-oxo-2-(tetradecylamino)ethyl] carbamate

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atmosphere for 90 mins. The TLC of the reaction mixture showed the formation of the acid was added. The mixture was stirred for another 35 mins, followed by the addition IH-tetrazole (319 mg, 4.56 mmol). To this mixture was added a 1:1 mixture of freshly distilled THF/CH2Cl2 (40 ml). After 10 mins, dibenzyldiisopropyl phosphoramidate of Na-metabisulfite to quench the excess peracetic acid. The THF and CH2Cl2 were To the pyridine-washed starting 27 (305 mg, 0.761 mmol) was added product. This mixture was cooled to 0 °C (ice bath), and a large excess of peracetic and was washed with Na-metabisulfite (2x30 ml), NaHCO₃ (2x40 ml), water (2x35 removed under reduced pressure. The concentrate was treated with EtOAc (70 ml), concentrated under reduced pressure. The residue was subjected to flash column (1.57 gm, 4.56 mmol) was added, and the reaction was stirred under an argon ml), and brine (2x35 ml). The organic portion was dried over NaSO4, and chromatography, eluting with EtOAchexanes of various compositions.

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2H), 3.16-3.23 (m, 2H), 4.02-4.09 (m, 1H), 4.31-4.43 (m, 2H), 5.00-5.15 (m, 4H), 5.57 Anal. Calcd. for C₃₆H₅₇N₂O₅P · 1H₂O · 0.5C₄H₅O₂: C, 63.14; H, 8.78; N, 3.88. Found: 80.57, 128.0, 128.05, 128.09, 128.58, 128.64, 128.68, 135.45, 135.54, 135.59, 168.51; vacuo to afford 451 mg (89 %) of 51 as a white waxy solid: mp 33-35°C; H NMR (CDCl₃) 8 0.87 (t, J=6.4 Hz, 3H), 1.23-1.25 (bm, 22H), 1.44 (s, 9H), 1.52-1.55 (m, Appropriate fractions were pooled, and concentrated to dryness in 22.67, 26.81, 28.27, 29.25, 29.33, 29.44, 29.51, 29.59, 29.62, 29.65, 31.91, 39.69, 46.49, 54.47, 67.00, 67.07, 67.24, 67.32, 69.66, 69.68, 69.74, 76.12, 77.20, 77.84, (bs, 1H), 6.34 (t, J=5.0 Hz, 1H) 7.31-7.40 (m, 10H); ¹³C (CDCl₃) 8 14.08, 19.03, C, 62.80; H, 8.38; N, 4.21.

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oxy)methyl}-2-(octadecylamino)-2-oxoethyl] tert-Butyl N-[1-{([dl(benzyloxy)phosphoryl] Compound 52:

atmosphere for 90 mins. The TLC of the reaction mixture showed the formation of the IH-tetrazole (329 mg, 4.70 mmol). To this mixture was added a 1:1 mixture of freshly acid was added. The mixture was stirred for another 35 mins, followed by the addition To the pyridine-washed starting 26 (270 mg, 0.783 mmol) was added distilled THF/CH2Cl2 (50 ml). After 10 mins, dibenzyldiisopropyl phosphoramidate of Na-metabisulfite to quench the excess peracetic acid. The THF and CH2Cl2 were product. This mixture was cooled to 0 °C (ice bath), and a large excess of peracetic and was washed with Na-metabisulfite (2x25 ml), NaHCO₃ (2x25 ml), water (2x25 removed under reduced pressure. The concentrate was treated with BtOAc (50 ml), concentrated under reduced pressure. The residue was subjected to flash column (1.62 gm, 4.70 mmol) was added, and the reaction was stirred under an argon ml), and brine (2x25 ml). The organic portion was dried over NaSO4, and chromatography, eluting with EtOAc/hexanes of various compositions.

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vacuo to afford 135 mg (28 %) of 52 as a white solid: mp 52-54°C; 'H NMR (CDC!₃) 2H), 4.01-4.09 (m, 1H), 4.30-4.44 (m, 2H), 5.00-5.05 (m, 4H), 5.56 (bs, 1H), 6.32 (bs, Calcd. for CapH63N2O,P · 0.75H2O · 1C,H8O2; C, 64.56; H, 9.17; N, 3.42. Found: C, 3349, 2919, 2852, 1717, 1685, 1654, 1516, 1242, 1163, 1037, 1025, 999 cm⁻¹; Anal. 5 0.87 (t, J=6.4 Hz, 3H), 1.23 (bm, 14H), 1.44 (s, 9H), 1.63 (m, 2H), 3.17-3.24 (m, 127.93, 128.06, 128.10, 128.65, 128.70, 128.73, 135.43, 168.51, 170.07; IR (KBr) 29.42, 29.52, 29.60, 29.64, 29.69, 31.91, 39.68, 67.00, 67.07, 69.69, 69.74, 77.20, Appropriate fractions were pooled, and concentrated to dryness in (H) 7.29-7.39 (m, 10H); ¹³C (CDCl₃) 8 14.11, 22.68, 26.80, 28.25, 29.26, 29.35, 54.23; H, 9.05; N, 3.78.

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Compound 53: ter-Butyl N-(1-{((di(benzyloxy)phosphoryl) oxy)methyl}-2-0xo-2-[4-(tetradecyloxy)anlilao)

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H-tetrazole (450 mg, 6.42 mmol). To this mixture was added a 1:1 mixture of freshly atmosphere for 90 mins. The TLC of the reaction mixture showed the formation of the ucid was added. The mixture was stirred for another 35 mins, followed by the addition To the pyridine-washed starting 29 (310 mg, 0.647 mmol) was added iistilled THF/CH2Cl2 (40 ml). After 10 mins, dibenzyldiisopropyl phosphoramidate product. This mixture was cooled to 0 °C (ice bath), and a large excess of peracetic of Na-metabisulfite to quench the excess peracetic acid. The THF and CH2Cl2 were (2.21 gm, 6.42 mmol) was added, and the reaction was stirred under an argon

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chromatography, cluting with BtOAc/hexanes of various compositions. concentrated under reduced pressure. The residue was subjected to flash column ml), and brine (2×35 ml). The organic portion was dried over NaSO4, and and was washed with Na-metabisulfite (2x25 ml), NaHCO₃ (2x35 ml), water (2x35 removed under reduced pressure. The concentrate was treated with EtOAc (70 ml),

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6.86 (m, J_0 =9.0 Hz, 2H) 7.31-7.39 (m, 12H), 8.82 (bs, 1H); 13 C (CDCl₃) δ 14.10, 3H), 4.01-4.16 (m, 1H), 4.42-4.49 (m, 2H), 4.96-5.09 (m, 4H), 5.65 (bs, 1H), 6.80-0.87 (t, J=6.5 Hz, 3H), 1.30 (s, 22H), 1.46 (s, 9H), 1.71-1.80 (m, 2H), 3.91 (t, J=6.5Hz, 2852, 1717, 1677, 1513, 1457, 1237, 1059, 998 cm⁻¹; Anal. Calcd. for C₄₂H₆₁N₂O₈P 114.79, 121.72, 128.07, 128.13, 128.65, 128.74, 130.03, 166.71; IR (KBr) 3340, 2920, 22.67, 26.02, 28.26, 29.26, 29.34, 29.40, 29.57, 29.64, 31.91, 68.31, 69.84, 77.20, vacuo to afford 81 mg (17 %) of 53 as a white solid: mp 74-76°C; ¹H NMR (CDCl₃) 8 1H₂O · 0.45C₆H₁₄: C, 66.31; H, 8.63; N, 3.46. Found: C, 65.92; H, 9.02; N, 3.84. Appropriate fractions were pooled, and concentrated to dryness in

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Compound 54: \(\alpha -\text{T-Butyl N-[1-\(([di(henzyloxy)phosphoryl]oxy)\)}\) methyl\rangle -\((-\text{methoxyanilino}\rangle -\text{oxoethyl}\)

column chromatography, eluting with EtOAc/hexanes of various compositions. NaSO4, and concentrated under reduced pressure. The residue was subjected to flash (2×25 ml), water (2×25 ml), and brine (2×25 ml). The organic portion was dried over with EtOAc (50 ml), and was washed with Na-metabisulfite (2×15 ml), NaHCO₃ excess of peracetic acid was added. The mixture was stirred for another 35 mins, THF and CH_2Cl_2 were removed on a rotary evaporator. The concentrate was treated followed by the addition of Na-metabisulfite to quench the excess peracetic acid. The formation of the product. This mixture was cooled to 0 °C (ice bath), and a large under an argon atmosphere for 90 mins. The TLC of the reaction mixture showed the phosphoramidate (1.25 gm, 3.625 mmol) was added, and the reaction was stirred freshly distilled THF/CH2Cl2 (20 ml). After 10 mins, dibenzyldiisopropyl 1H-tetrazole (254 mg, 3.625 mmol). To this mixture was added a 1:1 mixture of To the pyridine-washed starting 30 (225 mg, 0.725 mmol) was added

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5.72 (bs, 1H), 6.78-6.82 (m, J₆=9.0 Hz, 2H) 7.26-7.33 (m, 10H), 7.36-7.41 (m, J₆=9.0 8 1.44 (a, 9H), 4.11 (s, 3H), 4.09-4.18 (m, 1H), 4.43-4.51 (m, 2H), 4.98-5.05 (m, 4H), 135.42, 156.62, 166.75; ³¹P NMR (CDCl₃) & 16.72 (1P); IR (KB_r) 3337, 2969, 1716, Hz, 2H), 8.41 (bs, 1H); ¹³C (CDCl₃) 8 28.26, 55.45, 66.93, 67.00, 69.76, 69.83, 69.90, vacuo to afford 195 mg (47 %) of 54 as a white solid: mp 82-84°C; 'H NMR (CDCl₃) 77.20, 80.91, 114.11, 121.75, 128.06, 128.12, 128.64, 128.72, 128.73, 130.38, 135.28 Appropriate fractions were pooled, and concentrated to dryness in

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WO 01/71022

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PCT/US01/08729

61.05; H, 6.18; N, 4.91. Found: C, 60.80; H, 6.20; N, 4.88. 1689, 1665, 1514, 1457, 1304, 1245, 999 cm⁻¹; Anal. Calcd. for C₁₉H₃₅N₂O₆P: C,

Example 5 - Synthesis of Compounds 55-59

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Compound 55: 2-Amino-3-(nonylamino)-3-oxopropyl dihydrogen phosphate

Calcd. for C13H29N2O5P-0.5CH3OH: C, 47.64; H, 9.18; N, 8.23. Found: C, 47.24; H, 30.46, 30.94, 31.16, 31.30, 31.39, 33.81, 43.53, 57.21, 66.42, 167.86; MS m/z 323 (M-8 0.81-0.82 (m, 3H), 1.26-1.30 (m, 14H), 1.59 (m, 2H), 3.37-3.38 (m, 2H), 4.54-4.59 afford 48 mg (90 %) of 55 as a white powder: mp 196-198 °C; HNMR (CF3COOD) 8.84; N, 8.02. H); IR (KBr) 3314, 2920, 2853, 1670, 1575, 1477, 1246, 1063, 1043 cm⁻¹; Anal. (m, 1H), 4.72-4.81 (m, 2H); ¹³C NMR (CF₃COOD) & 14.66, 24.39, 28.60, 28.60, After 4 hours TLC determined the completion of the reaction, the reaction mixture was filtered through celite, and the cluste was concentrated under reduced pressure to 10 %Pd/C (catalytic amount). Hydrogenation was carried out for 4 hrs at 50 psi. To a solution of 50 (100 mg, 0.165 mmol) in EtOH (15 ml) was added

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Compound 56: 2-Amino-3-oxo-3-(tetradecylamino)propyl dihydrogen phosphate

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MS m/z 379 (M-H); IR (KBr) 3318, 2923, 2852, 1671, 1657, 1563, 1475, 1242, 1055 cm"; Anal. Calcd. for C₁₇H₃₇N₃O₅P: C, 53.67; H, 9.80; N, 7.36. Found: C, 53.40; H 31.22, 31.27, 33.62, 43.27, 56.96, 66.16, 167.60; ³¹P NMR (CE₃COOD) 8 17.93 (1P); 4.78 (m, 2H); 13C NMR (CF₃COOD) 8 14.43, 24.16, 28.34, 30.21, 30.69, 31.01, 31.17, afford 75 mg (90 %) of 56 as a white powder: mp 189-190 °C; ¹H NMR (CF₃COOD) filtered through celite, and the cluate was concentrated under reduced pressure to 8 0.81 (bs, 3H), 1.24 (s, 23H), 1.57 (m, 2H), 3.37 (m, 2H), 4.54-4.58 (m, 1H), 4.73-After 3 hours TLC determined the completion of the reaction, the reaction mixture was 10 %Pd/C (catalytic amount). Hydrogenation was carried out for 3 hrs at 45 psi. To a solution of \$1 (145 mg, 0.219 mmol) in EtOH (15 ml) was added

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Compound 56a: 2-(Acetylamino)-3-oxo-3-(tetradecylamino) propyl dihydrogen phosphate

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mixture was stirred with 20 ml of aqueous HCl. The acidic mixture was extracted Excess pyridine and acetic anhydride were on a rotary evaporator. The resultant large excess of acetic anhydride. The mixture was allowed to stir at r.t. overnight. To a sample of 56 (20 mg, 0.052 mmol) in 0.5 ml pyridine was added a

PCT/US01/08729

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3.20 (m, 2H), 4.10-4.28 (m, 2H), 4.54-4.62 (m, 1H); 13C NMR (CDC1,/CD3OD) 13.48, with BtOAc (2x25 ml). The BtOAc layer was washed with water (2x25 ml) and brine concentrated under reduced pressure to afford 15 mg (71%) of 56a as a gummy solid: H NNAR (CD5OD), 8 0.89 (t, J=6.3 Hz, 3H), 1.27 (s, 22H), 1.99-2.02 (m, 3H), 3.15-(2x25 ml). The organic portion was dried over NaSO4 and filtered. The eluate was 43.27, 56.96, 66.16, 163.02, 174.96; IR (KBr) 3316, 2923, 2853, 1671, 1657, 1560, 16.19, 22.23, 26.50, 28.91, 29.21, 31.48, 30.21, 31.01, 31.17, 31.22, 31.27, 33.62, 1467, 1247, 1059 cm⁻¹.

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Compound 57: 2-Amino-3-(octadecylamino)-3-oxopropyi dihydrogen phosphate

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After 4 hours TLC determined the completion of the reaction, the reaction mixture was 5 0.81 (t, 1=6.9 Hz, 3H), 1.25 (s, 31H), 1.58 (m, 2H), 3.34-3.44 (m, 2H), 4.49-4.59 (m, fford 70 mg (98 %) of 57 as a white powder: mp 190-192 °C; ¹H NMR (CF3COOD) To a solution of **52** (117 mg, 0.164 mmol) in EtOH (15 ml) was added 31.28, 31.31, 31.44, 31.48, 31.55, 33.89, 43.53, 57.12, 57.21, 66.35, 167.85; MS m/z Calcd. for C21H4sN2O5P: C, 57.77; H, 10.39; N, 6.42. Found: C, 57.61; H, 10.22; N, IH), 4.71-4.81 (m, 2H); ¹³C NMR (CF₃COOD) 5 14.70, 24.43, 28.60, 30.46, 30.95, 435 (M-H); IR (KBr) 3325, 2922, 2852, 1674, 1655, 1560, 1472, 1045 cm⁻¹; Anal. filtered through celite, and the cluate was concentrated under reduced pressure to 10 %Pd/C (catalytic amount). Hydrogenation was carried out for 4 hrs at 50 psi.

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Compound 58: 2-Amino-3-oxo-3-[4-(tetradecyloxy)anilino] propyl dihydrogen phosphate

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After 4 hours TLC determined the completion of the reaction, the reaction mixture was afford 22 mg (88 %) of 58 as a white powder: mp 187-190 °C; HNMR (CF3COOD) To a solution of \$3 (40 mg, 0.054 mmol) in EtOH (15 ml) was added 5 0.80-0.82 (m, 3H), 1.25 (m, 20H), 1.77-1.84 (m, 2H), 4.20 (t, 1=6.0 Hz, 2H), 4.64-4.74 (m, 1H), 4.90-4.91 (m, 2H), 7.04-7.07 (d, J₀=9.0 Hz, 2H), 7.32-7.35 (d, J₀=9.0 1240, 1046 cm.'; Anal. Calcd. for CzzH, NzOsP-0.5CH3OH-0.5CHCl3: C, 52.58; H, Hz, 2H); ¹³C NMR (CF₃C00D) 8 14.81, 24.54, 27.57, 30.62, 31.19, 31.38, 31.46, 167.06; MS m/z 471 (M-H); IR (KBr) 3325, 2923, 2852, 1665, 1553, 1515, 1469, filtered through celite, and the eluate was concentrated under reduced pressure to 10 %Pd/C (catalytic amount). Hydrogenation was carried out for 4 hrs at 50 psi. 31.52, 31.60, 31.65, 33.99, 57.70, 66.53, 73.66, 119.32, 126.55, 131.25, 158.87, 3.00; N, 5.11. Found: C, 52.89; H, 7.83; N, 5.29.

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2-Amino-3-(4-methoxyanilino)-3-oxopropyl dihydrogen phosphate Compound 59:

After 2 hours TLC determined the completion of the reaction, the reaction mixture was afford 82 ing (96 %) of 59 as a white powder: mp 199-202 °C; ¹H NMR (CF₃COOD) 1565, 1515, 1478, 1236, 1045 cm-1; Anal. Calcd. for CooH13N2O6P: C, 41.39; H, 5.21; To a solution of \$4 (125 mg, 0.219 mmol) in BtOH (15 ml) was added 8 3.93 (s, 3H), 4.65-4.75 (m, 1H), 4.88-4.94 (m, 2H), 7.01-7.04 (d, 10=9.0 Hz, 2H), filtered through celite, and the cluate was concentrated under reduced pressure to 126.64, 131.07, 159.62, 167.07; MS m/z 289 (M-H); IR (KBr) 3317, 2961, 1680, 10 %Pd/C (catalytic amount). Hydrogenation was carried out for 2 hrs at 45 psi. 7.31-7.34 (d, Jo=9.0 Hz, 2H); ¹³C NMR (CDCI₃) 8 57.60, 58.00, 66.54, 117.69, N, 9.65. Found: C, 41.25; H, 5.35; N, 9.73.

Example 6 - Synthesis of Intermediate Compounds 63-65

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argon atmosphere. The starting alcohol was washed with anhydrous pyridine (3 times) The glassware used was flame-dried and cooled to room temperature under an and dried on high vacuum for 48 hrs. The reaction was carried out in an argon atmosphere. THF and CH2Cl2 were freshly distilled prior to their use.

Compound 63: 1,2-(3-Octadecyloxypropane)-bis(dibenzylphosphate)

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nmol) was added 1H-tetrazole (229 mg, 3.26 mmol). To this mixture was added a 1:1 phosphoramidate (1.12 gm, 3.26 mmol) was added, and the reaction was stirred under mixture of freshly distilled THF/CH2Cl3 (50 ml). After 10 mins, dibenzyldiisopropyl followed by the addition of Na-metabisulfite to quench the excess peracetic acid. The (2×30 ml), water (2×30 ml), and brine (2×30 ml). The organic portion was dried over THF and CH₂Cl₂ were removed under reduced pressure. The concentrate was treated NaSO4, and concentrated under reduced pressure. The residue was subjected to flash To the pyridine-washed starting all-batyl alcohol (60, 225 mg, 0.652 formation of the product. This mixture was cooled to 0 °C (ice bath), and a large with EtOAc (70 ml), and was washed with Na-metabisulfite (2x25 ml), NaHCO3 excess of peracetic acid was added. The mixture was stirred for another 35 mins, an argon atmosphere for 90 mins. The TLC of the reaction mixture showed the column chromatography, eluting with EtOAc/hexanes of various compositions.

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vacuo to afford 303 mg (53 %) of 63 as a clear oil: ¹HNMR (CDCl3) 8 0.86 (t, 1=6.4 Hz, 3H), 1.24 (bm, 28H), 1.33-1.35 (m, 2H), 1.45 (m, 2H), 3.29-3.36 (m, 2H), 3.48-Appropriate fractions were pooled, and concentrated to dryness in

PCT/US01/08729

3.50 (d, J=5.2 Hz, 2H), 4.04-4.22 (m, 2H), 4.60 (m, 1H), 5.00 (m, 8H), 7.27-7.33 (m 20H); ¹³C (CDCl₃) 8 14.05, 18.96, 22.62, 25.95, 29.29, 29.41, 29.49, 29.53, 29.59, 29.63, 31.85, 46.48, 66.58, 69.20, 69.23, 69.28, 69.36, 71.75, 75.37, 127.76, 127.82, 127.86, 127.88, 127.94, 128.36, 128.45, 128.49, 128.61, 128.62, 135.46, 135.54, 135.59, 135.65, 135.68, 135.75, 135.79; MS m/z 866 (M+H)[†].

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Compound 64: 1,2-(3-Dodecyloxypropane) bb(dibenzylphosphate)

To the pyridine-washed starting dl-3-O-n-dodecyl-1,2-propanediol (61, 400 mg, 1.5 mmol) was added 1H-tetrazole (645 mg, 9.2 mmol). To this mixture was added a 1:1 mixture of freshly distilled THF/CH₂Ch₂ (40 ml). After 10 mins, dibenzyldiisopropyl phosphoramidate (3.18 gm, 9.2 mmol) was added, and the reaction was stirred under an argon atmosphere for 90 mins. The TLC of the reaction mixture showed the formation of the product. This mixture was cooled to 0 °C (ice bath), and a large excess of peracetic acid was added. The mixture was stirred for another 35 mins, followed by the addition of Na-metabisulfite to quench the excess peracetic acid. The THF and CH₂Cl₂ were removed under reduced pressure. The concentrate was treated with EtOAc (80 ml), and was washed with Na-metabisulfite (2x35 ml), NaHCO₃ (2x40 ml), water (2x30 ml), and brine (2x30 ml). The organic portion was dried over NaSO₄, and concentrated under reduced pressure. The residue was subjected to flash column chromatography, eluting with EtOAc/hexanes of various compositions.

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Appropriate fractions were pooled, and concentrated to dryness in vacuo to afford 100 mg (< 10 %) of 64 as a clear oil: ¹H NMR (CDCl₃) 8 0.86 (t, J=6.3 Hz, 3H), 1.23 (bm, 18H), 1.46 (m, 2H), 3.13-3.36 (m, 2H), 3.49-3.51 (d, J=5.2 Hz, 2H), 4.03-4.23 (m, 2H), 4.59 (m, 1H), 5.01 (m, 8H), 7.26-7.34 (m, 20H); ¹¹C (CDCl₃) 8 14.11, 22.68, 26.01, 29.35, 29.47, 29.54, 29.59, 29.63, 29.66, 31.91, 69.01, 69.06, 69.26, 69.30, 69.34, 69.42, 69.62, 71.83, 77.21, 127.83, 127.89, 127.94, 127.95, 128.44, 128.52, 128.56, 135.64, 135.74, 135.85; IR (NaCl, neat) 3427, 1276, 1000, 885, 499 cm²; MS m/z 781 (M+H) †, m/z 803 (M+Na)†.

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Compound 65: 1,2-(3-Hexadecyloxypropane) bis(dibenzylphosphate)

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To the pyridine-washed starting dl-3-O-n-hexadecyl-1,2-propanediol (62, 500 mg, 1.57 mmol) was added 1H-tetrazole (664 mg, 9.47 mmol). To this mixture was added a 1:1 mixture of freshly distilled THF/CH₂Cl₂ (50 ml). After 10 mins, dibenzyldiisopropyl phosphoramidate (3.27 gm, 9.47 mmol) was added, and the reaction was stirred under an argon atmosphere for 90 mins. The TLC of the reaction

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WO 01/71022 - 57 -

PCT/US01/08729

mixture showed the formation of the product. This mixture was cooled to 0 °C (ice bath), and a large excess of peracetic acid was added. The mixture was stirred for another 35 mins, followed by the addition of Na-metabisulfite to quench the excess peracetic acid. The THF and CH₂Cl₂ were removed under reduced pressure. The concentrate was treated with EtOAc (80 ml), and was washed with Na-metabisulfite (2×35 ml), NaHCO₃ (2×40 ml), water (2×30 ml), and brine (2×30 ml). The organic portion was dried over NaSO₄, and concentrated under reduced pressure. The residue was subjected to flash column chromatography, eluting with BtOAc/hexanes of various compositions.

Appropriate fractions were pooled, and concentrated to dryness in vacuo to afford 205 mg (15 %) of 65 as a clear oil: 'H NMR (CDCl₃) \(\delta \) 0.87 (t, J=6.3 Hz, 3H), 1.25 (bm, 26H), 1.46 (m, 2H), 3.30-3.42 (m, 2H), 3.49-3.51 (d, J=5.2 Hz, 2H), 3.97-4.23 (m, 2H), 4.60 (m, 1H), 5.01 (m, 8H), 7.26-7.35 (m, 20H); \(^{13}\)C (CDCl₃) \(\delta \) 14.11, 22.68, 26.00, 29.35, 29.47, 29.54, 29.59, 29.64, 29.68, 31.91, 69.00, 69.06, 69.26, 69.29, 69.34, 69.41, 71.82, 71.74,75.52, 75.60, 77.20, 126.97, 127.82, 127.88, 127.93, 127.95, 127.99, 128.43, 128.51, 128.55, 128.60, 135.63, 135.73, 135.79, 135.83; IR (NaCl, neat) 3423, 1269, 1016, 736, cm⁻¹; MS m/z 837 (M+H) \(^+\), m/z 859 (M+Na)\(^+\).

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Example 7 - Synthesis of Compounds 66-68

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Compound 66: 1,2-(3-Octadecyloxypropane)-bis(dihydrogen phosphate)

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To a solution of 63 (135 mg, 0.156 mmol) in EtOH (15 ml) was added 10 %Pd/C (catalytic amount). Hydrogenation was carried out for 4 hrs at 60 psi. After 4 hours, TLC determined the completion of the reaction, the reaction mixture was filtered through celite, and the cluate was concentrated under reduced pressure to afford 70 mg (89 %) of 66 as a clear wax: H NMR (CD₃OD) & 0.89 (t, J=6.4 Hz, 3H), 1.28 (s, 30H), 1.55 (m, 2H), 3.45-3.50 (m, 2H), 3.62-3.64 (m, 2H), 4.00-4.16 (m, 2H), 4.47 (m, 1H); ¹⁹C NMR (CD₃OD) & 14.43, 19.30, 23.73, 27.20, 30.47, 30.64, 30.78, 33.07, 72.80; MS m/z 503 (M-H); IR (NaCl Next) 1011 cm⁻¹.

Compound 67: 1,2-(3-Dodecyloxypropane)-bls(dihydrogen phosphate)

To a solution of 64 (70 mg, 0.089 mmol) in EtOH (15 ml) was added 10 %Pd/C (catalytic amount). Hydrogenation was carried out for 4 hrs at 60 psi.

After 4 hours, TLC determined the completion of the reaction, the reaction mixture was filtered through celite, and the cluste was concentrated under reduced pressure to

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- 58 -

afford 35 mg (94 %) of 67 as a clear wax. ¹H NMR (CD₂OD) 8 0.79 (t, J=6.7 Hz, 3H), 1.90 (s, 18H), 1.46 (m, 2H), 3.34-3.41 (m, 2H), 3.49-3.73 (m, 2H), 3.78-4.05 (m, 2H), 4.47 (m, 1H); ¹³C NMR (CD₂OD) 5 14.43, 23.71, 23.74, 27.20, 30.49, 30.64, 30.76, 30.81, 33.08, 66.80, 72.79, MS m/z 419 (M-H); IR (NaCl Neat) 1008 cm⁻¹.

Compound 68: 1,2-(3-Hexadecyloxypropaue)-bis(dihydrogen phosphate)

To a solution of 65 (138 mg, 0.164 mmol) in EtOH (15 mJ) was added 10 %Pd/C (catalytic amount). Hydrogenation was carried out for 4 hrs at 60 psi.

After 4 hours, TLC determined the completion of the reaction, the reaction mixture was filtered through celite, and the cluate was concentrated under reduced pressure to afford 75 mg (96 %) of 68 as a clear wax: ¹H NMR (CD₃OD) 8 0.89 (t. J=6.4 Hz, 3H), 1.28 (s, 23H), 1.56 (m, 2H), 3.43-3.50 (m, 2H), 3.58-3.65 (m, 2H), 3.89-4.16 (m, 2H), 4.47 (m, 1H); ¹³C NMR (CD₃OD) 8 14.44, 23.74, 27.20, 30.48, 30.64, 30.80, 33.08, 72.80; MS m/z 475 (M-H); IR (NaCl Neat) 1011 cm⁻¹

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Example 8 - Synthesis of Intermediate Compounds 77-84

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The glassware used was flame-thied and cooled to room temperature under an argon atmosphere. The starting alcohol was washed with anhydrous pyridine (3 times) and dried on high vacuum for 48 hrs. The reaction was carried out in an argon atmosphere. THF and CH₂Cl₂ were freshly distilled prior to their use.

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Compound 77: 1,2-(3-Tetradecanoyloxypropane)bis(dibenzylphosphate)

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To the pyridine-washed starting monomyristine (69, 800 mg, 2.6 mmol) was added 1H-tetrazole (1.01 gm, 14.5 mmol). To this mixture was added freshly distilled THF (45 ml). After 10 mins, dibenzy/diisopropyl phosphoramidate (5.02 gm, 14.5 mmol) was added, and the reaction was stirred under an argon atmosphere for 90 mins. The TLC of the reaction mixture showed the formation of the product. This mixture was cooled to 0 °C (ice bath), and a large excess of peracetic acid was added. The mixture was stirred for another 35 mins, followed by the addition of Namerabisulfite to quench the excess peracetic acid. The THF was removed under reduced pressure. The concentrate was treated with EtOAc (100 ml), and was washed with Na-menbisulfite (2×50 ml), NaHCO₃ (2×75 ml), water (2×50 ml), and brine (2×50 ml). The organic portion was dried over NaSO₄, and concentrated under reduced pressure. The residue was subjected to flash column chromatography, eluting with EtOAc/hexanes of various compositions.

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WO 01/71022

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PCT/US01/08729

Appropriate fractions were pooled, and concentrated to dryness in vacuo to afford 600 mg (28 %) of 77 as a clear oil: 'H NMR (CDCi₃) δ 0.87 (t, J=6.3 Hz, 31.25 (bm, 20H), 1.53 (m, 2H), 2.17-2.32 (m, 2H), 3.96-4.24 (m, 4H), 4.61-4.70 (m, 1H), 4.99-5.08 (m, 8H), 7.29-7.35 (m, 20H); ¹³C (CDCi₃) δ 14.10, 22.67, 24.70, 29.08, 29.23, 29.33, 29.44, 29.59, 29.62, 29.66, 31.90, 33.86, 64.24, 65.82, 69.41, 69.46, 69.48, 69.53, 69.57, 77.20, 127.85, 127.91, 127.98, 127.99, 128.04, 128.57, 128.59, 128.70, 128.59, 138.

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Compound 78: 1,2-(3-Pentadecanoyloxypropane)bis(dibenzylphosphate)

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To the pyridine-washed starting monopentadecanoin (70, 800 mg, 2.5 mmol) was added 1H-tetrazole (970 mg, 13.9 mmol). To this mixture was added freshly distilled THF (45 ml). After 10 mins, dibenzyldiisopropyl phosphoramidate (4.80 gm, 13.9 mmol) was added, and the reaction was stirred under an argon atmosphere for 90 mins. The TLC of the reaction mixture showed the formation of the product. This mixture was cooled to 0°C (ice bath), and a large excess of peracetic acid was added. The mixture was tirred for another 35 mins, followed by the addition of Na-metabisulfite to quench the excess peracetic acid. The THF was removed under reduced pressure. The concentrate was treated with EtOAc (100 ml), and was washed with Na-metabisulfite (2x50 ml), NaHCO₇ (2x100 ml), water (2x50 ml), and brine (2x50 ml). The organic portion was dried over NaSO₆, and concentrated under reduced pressure. The residue was subjected to flash column chromatography, eluting with EtOAcohexanes of various compositions.

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Appropriate fractions were pooled, and concentrated to dryness in vacuo to afford 741 mg (35 %) of 78 as a clear oil: ¹H NMR (CDCl₃) 8 0.87 (t, J=6.4 Hz, 3H), 1.25 (bm, 22H), 1.53 (m, 2H), 2.17-2.32 (m, 2H), 3.95-4.24 (m, 4H), 4.61-4.70 (m, 1H), 4.99-5.07 (m, 8H), 7.29-7.35 (m, 20H); ¹³C (CDCl₃) 8 14.09, 22.66, 24.69, 29.03, 29.23, 29.33, 29.44, 29.59, 29.65, 21.89, 33.85, 64.23, 65.86, 69.40, 69.46, 69.48, 69.53, 69.56, 77.20, 127.84, 127.90, 127.97, 127.98, 128.03, 128.55, 128.69, 128.71 135.50, 135.59, 173.09; IR (NaCl, Next) 3421, 1742, 1457, 1275, 1035, 1014, 1001 cm⁻¹; MS m/z 837 (M+H)⁺, m/z 859 (M+Na)⁺.

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Compound 79: 1,2-(3-Hexadecanoyloxypropane)bis(dibenzylphosphate)

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To the pyridine-washed starting monopalmitin (71, 800 mg, 2.4 mmol) was added 1H-tetrazole (1.00 gm, 14.2 mmol). To this mixture was added freshly distilled THP (45 ml). After 10 mins, dibenzyldiisopropyl phosphoramidate (4.90 gm,

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14.2 mmol) was added, and the reaction was stirred under an argon atmosphere for 90 mins. The TLC of the reaction mixture showed the formation of the product. This mixture was cooled to 0 °C (ice bath), and a large excess of peracetic acid was added. The mixture was stirred for another 35 mins, followed by the addition of Nametabisulfite to quench the excess peracetic acid. The THF was removed under reduced pressure. The concentrate was treated with BtOAc (100 ml), and was washed with Na-metabisulfite (2×50 ml), NaHCO₃ (2×100 ml), water (2×50 ml), and brine (2×50 ml). The organic portion was dried over NaSO₄, and concentrated under reduced pressure. The residue was subjected to flash column chromatography, eluting with BtOAc/hexanes of various compositions.

Appropriate fractions were pooled, and concentrated to dryness in vacuo to afford 786 mg (38 %) of 79 as a clear oil: ¹H NMR (CDCl₃) \$ 0.87 (t, J=6.4 Hz, 3H), 1.25 (bm, 24H), 1.53 (m, 2H), 2.17-2.32 (m, 2H), 3.96-4.24 (m, 4H), 4.61-4.70 (m, 1H), 4.99-5.08 (m, 8H), 7.29-7.35 (m, 20H); ¹¹C (CDCl₃) \$ 14.09, 22.66, 24.71, 29.09, 29.23, 29.33, 29.45, 29.60, 29.63, 29.67, 31.90, 33.87, 62.23, 62.30, 65.89, 69.43, 69.48, 69.50, 69.55, 69.58, 77.20, 126.96, 127.85, 127.91, 127.98, 128.04, 128.56, 128.59, 128.64, 128.71 135.52, 135.61, 173.07; IR (NaCl, Neat) 3421 1742, 1457, 1273, 1035, 1016, 1001 cm⁻¹; MS m/z 851 (M+H)⁺, m/z 873 (M+Na)⁺.

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Compound 80: 1,2-(3-Heptadecanoyloxypropane)bis(dibenzylphosphate)

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To the pyridine-washed starting monoheptadecanoin (72, 800 mg, 2.32 mmol) was added 1H-tetrazole (980 mg, 13.9 mmol). To this mixture was added freshly distilled THF (40 ml). After 10 mins, dibenzyldiisopropyl phosphoramidate (4.81 gm, 13.9 mmol) was added, and the reaction was stirred under an argon atmosphere for 90 mins. The TLC of the reaction mixture showed the formation of the product. This mixture was cooled to 0 °C (ice bath), and a large excess of peracetic acid was added. The mixture was stirred for another 35 mins, followed by the addition of Na-metabisulfite to quench the excess peracetic acid. The THF was removed under reduced pressure. The concentrate was treated with EiOAc (100 ml), and was washed with Na-metabisulfite (2×50 ml), NaHCO₃ (2×100 ml), water (2×50 ml), and brine (2×50 ml). The organic portion was dried over NaSO₄, and concentrated under reduced pressure. The residue was subjected to flash column chromatography, eluting with EiOAc/hexanes of various compositions.

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Appropriate fractions were pooled, and concentrated to dryness in vacuo to afford 1.48 gm (74 %) of 80 as a clear oil: ¹H NMR (CDCl₃) 8 0.87 (t, 1=6.4 Hz, 3H), 1.23-1.25 (bm, 26H), 1.53 (m, 2H), 2.20 (t, 1=7.1 Hz, 2H), 4.02-4.24 (m, 4H), 4.66 (m, 1H), 4.99-5.05 (m, 8H), 7.29-7.35 (m, 20H); ¹³C (CDCl₃) 8 14.10,

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PCT/USII1/IB729

22.66, 24.69, 29.07, 29.23, 29.33, 29.44, 29.59, 29.63, 29.66, 31.89, 33.84, 62.21, 62.27, 65.85, 69.40, 69.45, 69.47, 69.52, 69.56, 74.04, 74.23, 77.20, 127.83, 127.87, 127.96, 127.97, 128.53, 128.55, 128.57, 128.59, 135.47, 135.56, 173.07; IR (NaCl, Neat) 3483, 1743, 1457, 1281, 1035, 1013, 1000 cm⁻¹; MS m/z 865 (M+H) ⁺, m/z 887 (M+Na)⁺.

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Compound 81: 1,2-(3-Octudecanoyloxypropane) bis(dibenzylphosphate)

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was added 1H-tetrazole (1.00 gm, 14.2 mmol). To this mixture was added freshly distilled THF (40 ml). After 10 mins, dibenzyldiisopropyl phosphoramidate (4.92 gm, 14.2 mmol) was added, and the reaction was stirred under an argon atmosphere for 90 mins. The TLC of the reaction mixture showed the formation of the product. This mixture was cooled to 0 °C (ice bath), and a large excess of peracetic acid was added. The mixture was stirred for another 35 mins, followed by the addition of Namethisulfite to quench the excess peracetic acid. The THF was removed under reduced pressure. The concentrate was treated with BtOAc (100 ml), and was washed with Na-metabisulfite (2×50 ml), NaHCO₃ (2×100 ml), water (2×50 ml), and brine (2×50 ml). The organic portion was dried over NaSO₄, and concentrated under reduced pressure. The residue was subjected to flash column chromatography, cluting with EtOAc/hexanes of various compositions.

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Appropriate fractions were pooled, and concentrated to dryness in vacuo to afford 870 mg (45 %) of 81 as a clear oil: 'H NMR (CDCl₃) & 0.87 (t, J=6.4 Hz, 3H), 1.23-1.25 (bm, 28H), 1.53 (m, 2H), 2.20 (t, J=7.2 Hz, 2H), 3.97-4.24 (m, 4H), 4.66 (m, 1H), 4.99-5.07 (m, 8H), 7.29-7.35 (m, 20H); ¹³C (CDCl₃) & 14.09, 22.66, 24.69, 29.08, 29.23, 29.33, 29.45, 29.59, 29.63, 29.67, 31.89, 33.85, 62.22, 62.28, 64.23, 65.87, 68.69, 69.23; 69.42, 69.50, 69.54, 69.58, 74.07, 74.25, 127.60, 127.84, 127.90, 127.98, 128.03, 128.54, 128.56, 128.58, 128.60, 128.71, 135.47, 135.57, 173.08; IR (NaCl, Next) 3421, 1742, 1457, 1273, 1251, 1216, 1035, 1016, 1000 cm⁻¹; MS m/z 879 (M+H) ⁺, m/z 991 (M+Na) ⁺.

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Compound 82: 1,2-(3-Nonadecanoyloxypropane)bis(dibenzylphosphate)

To the pyridine-washed starting Monononadecanoin (74, 800 mg, 2.1 mmol) was added 1H-ternazole (977 gm, 13.9 mmol). To this mixture was added freshly distilled THF (40 ml). After 10 mins, dibenzyldiisopropyl phosphoramidate (4.81 gm, 13.9 mmol) was added, and the reaction was stirred under an argon atmosphere for 90 mins. The TLC of the reaction mixture showed the formation of the

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of Na-metabisulfite to quench the excess peracetic acid. The THP was removed under acid was added. The mixture was stirred for another 35 mins, followed by the addition reduced pressure. The residue was subjected to flash column chromatography, eluting reduced pressure. The concentrate was treated with BtOAc (100 ml), and was washed product. This mixture was cooled to 0 °C (ice bath), and a large excess of peracetic with Na-metabisulfite (2x50 ml), NaHCO₃ (2x125 ml), water (2x75 ml), and brine (2×50 ml). The organic portion was dried over NaSO4, and concentrated under with EtOAc/hexanes of various compositions.

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vacuo to afford 1.47 gm (78 %) of 82 as a clear oil: ¹H NMR (CDCl₃) 8 0.87 (t, J=6.3 65.84, 69.38, 69.46, 69.51, 69.54, 74.03, 74.10, 74.15, 74.22, 77.20, 127.82, 127.88, 127.96, 128.53, 128.56, 135.45, 135.55, 173.06; IR (NaCl, Neat) 3483, 1743, 1457, Hz; 3H), 1.23-1.25 (bm, 30H), 1.53 (m, 2H), 2.20 (t, J=7.2 Hz, 2H), 4.02-4.24 (m, 22.65, 24.67, 29.06, 29.22, 29.32, 29.43, 29.58, 29.61, 29.66, 31.88, 33.83, 62.25, Appropriate fractions were pooled, and concentrated to dryness in 4H), 4.66 (m, 1H), 4.99-5.03 (m, 8H), 7.29-7.36 (m, 20H); ¹³C (CDCl₃) § 14.08, 1273, 1282, 1216, 1035, 1013 cm⁻¹; MS m/z 893 (M+H) ⁺, m/z 915 (M+Na)⁺.

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Compound 83: 1,2-(3-icosanoyloxypropane)-bis(dibenzylphosphate)

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etmosphere for 90 mins. The TLC of the reaction mixture showed the formation of the acid was added. The mixture was stirred for another 35 mins, followed by the addition of Na-metabisulfite to quench the excess peracetic acid. The THF was removed under reduced pressure. The concentrate was treated with EtOAc (100 ml), and was washed reduced pressure. The residue was subjected to flash column chromatography, eluting freshly distilled THF (40 ml). After 10 mins, dibenzyldiisopropyl phosphoramidate product. This mixture was cooled to 0 °C (ice bath), and a large excess of peracetic with Na-metabisulfite (2x50 ml), NaHCO₃ (2x125 ml), water (2x75 ml), and brine mmol) was added 1H-tetrazole (1.00 gm, 14.2 mmol). To this mixture was added To the pyridine-washed starting Monoarachidin (75, 800 mg, 2.06 (2×50 ml). The organic portion was dried over NaSO4, and concentrated under (4.92 gm, 14.2 mmol) was added, and the reaction was stirred under an argon with BtOAc/hexanes of various compositions.

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vacuo to afford 1.39 gm (74 %) of 83 as a clear oil: ¹H NNAR (CDCl₃) § 0.87 (t, J=6.4 Hz, 3H), 1.23-1.25 (bm, 32H), 1.53 (m, 2H), 2.20 (t, J=7.2 Hz, 2H), 4.02-4.24 (m, Appropriate fractions were pooled, and concentrated to dryness in 62.27, 65.86, 69.40, 69.45, 69.48, 69.52, 69.56, 74.05, 74.12, 74.16, 74.24, 77.20, 22.65, 24.69, 29.07, 29.23, 29.33, 29.44, 29.59, 29.63, 29.67, 31.89, 33.84, 62.21, 4H), 4.66 (m, 1H), 4.99-5.05 (m, 8H), 7.29-7.36 (m, 20H); ¹³C (CDCl₃) δ 14.09,

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WO 01/71022

PCT/US01/08729 ŝ 127.83, 127.89, 127.97, 128.53, 128.55, 128.57, 128.59, 135.47, 135.56, 173.07; 田 (NaCl, Neat) 3483, 1743, 1457, 1273, 1282, 1216, 1035, 1012, 1000 cm-1; MS m/z 907 (M+H)⁺, m/z 929 (M+Na)⁺.

Compound 84: 1,2-(3-Docosanoyloxypropane)-bis(dibenzylphosphate)

To the pyridine-washed starting Monobehenin (76, 800 mg, 1.92 mmol) distilled THF (40 ml). After 10 mins, dibenzyldiisopropyl phosphoramidate (5.14 gm, 14.8 mmol) was added, and the reaction was stirred under an argon atmosphere for 90 educed pressure. The residue was subjected to flash column chromatography, cluting reduced pressure. The concentrate was treated with BtOAc (100 ml), and was washed mixture was cooled to 0 °C (ice bath), and a large excess of peracetic acid was added mins. The TLC of the reaction mixture showed the formation of the product. This with Na-metabisulfite (2x50 ml), NaHCO3 (2x125 ml), water (2x75 ml), and brine was added 1H-tetrazole (1.00 gm, 14.2 mmol). To this mixture was added freshly netabisulfite to quench the excess peracetic acid. The THF was removed under (2x50 ml). The organic portion was dried over NaSO4, and concentrated under The mixture was stirred for another 35 mins, followed by the addition of Nawith EtOAc/hexanes of various compositions.

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Hz, 2H), 4.02-4.24 (m, 4H), 4.66 (m, 1H), 4.99-5.03 (m, 8H), 7.29-7.36 (m, 20H); ¹³C (CDCl₃) 8 14.08, 22.65, 24.68, 29.07, 29.22, 29.32, 29.44, 29.59, 29.62, 29.66, 31.88, 33.84, 62.20, 62.26, 65.85, 69.40, 69.45, 69.48, 69.53, 69.57, 74.05, 74.16,m 74.24, (CDCl₃) 8 0.87 (t, 1=6.4 Hz, 3H), 1.23-1.25 (bm, 36H), 1.53 (m, 2H), 2.20 (t, 1=7.2 Appropriate fractions were pooled, and concentrated to dryness in 77.20, 127.83, 127.88, 127.96, 127.97, 128.30, 128.52, 128.54, 128.57, 128.58, acuo to afford 1.27 gm (71 %) of 84 as a white wax like compound: 'H NMR 135.46, 135.55, 173.07; MS m/z 935 (M+H) *, m/z 957 (M+Na) *.

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Example 9 - Synthesis of Compounds 85-92

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Compound 85: 1,2-(3-Tetradecanoyloxypropane)-bis(dibydrogen phosphate)

was filtered through celite, and the cluate was concentrated under reduced pressure to)H), 1.28 (s, 20H), 1.56-1.63 (m, 2H), 2.24-2.38 (m, 2H), 3.93-4.42 (m, 4H), 4.59 (m, To a solution of 77 (385 mg, 0.468 mmol) in EtOH (15 ml) was added afford 210 mg (98 %) of 85 as a white wax: 1H NMR (CD30D) 8 0.89 (t, J=6.4 Hz, After 4 hours, TLC determined the completion of the reaction, the reaction mixture 10 %Pd/C (catalytic amount). Hydrogenation was carried out for 4 hrs at 60 psi.

- 64 - PCT/US01/08729

1H); ¹³C NMR (CD₃OD) \(\delta\) 14.44, 23.73, 26.09, 30.71, 30.23, 30.43, 30.47, 30.61, 30.75, 33.07, 34.80, 34.94, 61.90 61.96, 63.96, 63.70, 66.24, 74.33, 77.51, 175.02; MS m/z 461 (M-H); IR (NaCl Neat) 3386, 1702, 1216, 1019 cm⁻¹.

Compound 86: 1,2-(3-Pentadecanoyloxypropane)-bis(dihydrogen phosphate)

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To a solution of 78 (451 mg, 0.538 mmol) in EtOH (15 ml) was added 10 %Pd/C (catalytic amount). Hydrogenation was carried out for 4 hrs at 60 psi. After 4 hours, TLC determined the completion of the reaction, the reaction mixture was filtered through celite, and the cluate was concentrated under reduced pressure to afford 250 mg (97 %) of 86 as a white wax: ¹H NMR (CD₃OD) 8 0.89 (t, J=6.4 Hz, 3H), 1.28 (s, 22H), 1.58 (m, 2H), 2.24-2.38 (m, 2H), 3.97-4.21 (m, 4H), 4.38 (m, 1H); ¹³C NMR (CD₃OD) 8 14.44, 23.74, 26.05, 30.16, 30.36, 30.48, 30.57, 30.76, 33.08, 35.11, 61.36, 63.70, 63.90 66.24, 67.77, 70.22, 77.33, 77.40, 77.51, 175.63; MS m/z 475 (M-H); IR (NaCl Neat) 3380, 1728, 1216, 1031 cm⁻¹.

Compound 87: 1,2-(3-Hexadectanoyloxypropane)-bis(dihydrogen phosphate)

To a solution of 79 (561 mg, 0.659 mmol) in EtOH (15 ml) was added 10 %Pd/C (610 mg). Hydrogenation was carried out for 4 hrs at 60 psi. After 4 hours, TLC determined the completion of the reaction, the reaction mixture was filtered through celite, and the cluate was concentrated under reduced pressure to afford 300 mg (92 %) of 87 as a white was: ¹H NMR (CD₂OD) & 0.89 (t, J=6.4 Hz, 3H), 1.28 (s, 24H), 1.56-1.63 (m, 2H), 2.24-2.38 (m, 2H), 3.95-4.40 (m, 4H), 4.39 (m, 1H); ¹³C NMR (CD₂OD) & 14.43, 23.73, 25.89, 26.05, 26.09, 30.15, 30.23, 30.36, 30.44, 30.47, 30.56, 30.61, 30.67, 30.75, 33.07, 34.08, 34.94, 35.11, 61.36, 64.00, 66.22, 67.74, 70.22, 77.33, 77.40, 77.51, 175.03; MS m/z 489 (M-H); IR (NaCl Neat) 3357, 1729, 1216, 1029 cm⁻¹.

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Compound 88: 1,2-(3-Heptadecanoyloxypropane)-bis(dihydroger phosphate)

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To a solution of 80 (636 mg, 0.736 mmol) in EtOH (15 ml) was added 10 %Pd/C (724 mg). Hydrogenation was carried out for 4 hrs at 60 psi. After 4 hours, TLC determined the completion of the reaction, the reaction mixture was filtered through celite, and the cluate was concentrated under reduced pressure to afford 365 mg (98 %) of 88 as a white wax: ¹H NMR (CD₃OD) 8 0.89 (t, 1=6.6 Hz, 3H), 1.28 (s, 26H), 1.56-1.63 (m, 2H), 3.96-4.17 (m, 4H), 4.22-4.42 (m, 1H); ¹³C NMR (CD₃OD) 8 14.54, 23.73, 25.90, 26.10, 30.16, 30.24, 30.36, 30.43, 30.47, 30.56, 30.61, 30.76, 33.07, 34.81, 34.95, 61.37, 61.92, 63.97, 66.26, 67.70, 67.78, 70.06, 74.42, 77.46,

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WO 01/71022 - 65 -

PCT/US01/08729

175.04; MS m/z 503 (M-H); IR (NaCl Neat) 3357, 1710, 1216, 1032 cm⁻¹; Anal Calcd for C₂₀H₂₀O₁₀P₂ · 1H₂O: C, 45.97; H, 8.49. Found: C, 46.32; H, 8.73.

Compound 89: 1,2-(3-Octadecanoyloxypropane)-bis(dihydrogen phosphate)

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To a solution of 81 (530 mg, 0.603 mmol) in EtOH (15 ml) was added 10 %Pd/C (617 mg). Hydrogenation was carried out for 4 his at 60 psi. After 4 hours TLC determined the completion of the reaction, the reaction mixture was filtered through ceitie, and the eluste was concentrated under reduced pressure to afford 305 mg (97 %) of 89 as a white wax: HNMR (CD₂OD) 8 0.89 (t, j=6.3 Hz, 3H), 1.28 (s, 28H), 1.56-1.61 (m, 2H), 2.42-2.38 (m, 2H), 3.91-4.17 (m, 4H), 4.24-4.42 9m, 1H); ¹³C NMR (CD₂OD) 8 14.43, 23.74, 25.90, 26.06, 26.10, 30.16, 30.24, 30.36, 30.47, 30.57, 30.61, 30.67, 30.76, 33.08, 34.81, 34.95, 35.11, 61.37, 63.72, 66.26, 67.68, 67.75, 70.25, 77.48, 175.04; MS m/z 517 (M-H); IR (NaCl Neat) 3388, 1731, 1216, 1020 cm⁻¹.

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Compound 90: 1,2-(3-Nonadecanoyloxypropane)-bis(dihydrogen phosphate)

To a solution of 82 (952 mg, 1.06 mmol) in EtOH (25 ml) was added 10 %Pd/C (1.00 gm). Hydrogenation was carried out for 4 hrs at 60 psi. After 4 hours TLC determined the completion of the reaction, the reaction mixture was filtered through celite, and the eluate was concentrated under reduced pressure to afford 555 mg (98 %) of 90 as a white wax: HNMR (CD₂OD) 8 0.89 (t, J=6.4 Hz, 3H), 1.27 (s, 29H), 1.56-1.63 (m, 2H), 2.24-2.38 (m, 2H), 4.06-4.17 (m, 2H), 4.22-4.42 (m, 2H), 4.59 (m, 1H); ¹³C NMR (CD₂OD) 8 14.44, 23.74, 25.90, 26.06, 30.16, 30.24, 30.36, 30.48, 30.57, 30.63, 30.76, 30.79, 33.08, 34.81, 35.12, 63.94, 66.25, 175.03; MS m/z 531 (M-H); IR (NaCl Neat) 1735, 1216, 1012 cm⁻¹.

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Compound 91: 1,2-(3-Icosanoyloxypropane)-bis(dihydrogen phosphate)

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To a solution of 83 (711 mg, 0.784 mmol) in EtOH (25 ml) was added 10 %Pd/C (813 mg). Hydrogenation was carried out for 4 hrs at 60 psi. After 4 hours TLC determined the completion of the reaction, the reaction mixture was filtered through celitc, and the cluate was concentrated under reduced pressure to afford 419 mg (97 %) of 91 as a white wax: ¹H NMR (CD₂OD) δ 0.89 (t, 1=6.4 Hz, 3H₂, 1.28 (s, 32H), 1.58 (m, 2H), 2.24-2.38 (m, 2H), 3.95-4.42 (m, 4H), 4.58 (m, 1H); ¹¹C NMR (CD₂OD) δ 14.44, 23.74, 25.90, 26.06, 30.16, 30.24, 30.36, 30.48, 30.57, 30.63, 30.67, 33.08, 34.81, 35.11, 61.37, 61.98, 66.26, 67.69, 67.77, 77.42, 175.03; MS m/z 545 (M-H); IR (NaCl Neat) 3418, 1735, 1261, 1019 cm⁻¹.

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WO #1/71022

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PCT/US01/II8729

- 99 -

Compound 92: 1,2-(3-Docosanoyloxypropane)-bis(dihydrogen phosphate)

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To a solution of 84 (663 mg, 0.709 mmol) in E1OH (25 ml) was added 10 %Pd/C (710 mg). Hydrogenation was carried out for 4 hrs at 60 psi. After 4 hours TLC determined the completion of the reaction, the reaction mixture was filtered through celite, and the eluate was concentrated under reduced pressure to afford 400 mg (98 %) of 92 as a white wax: ¹H NMR (CD,OD) 8 0.89 (t, J=6.3 Hz, 3H), 1.27 (s, 36H), 1.58 (m, 2H), 2.24-2.38 (m, 2H), 3.98-4.42 (m, 4H), 4.59 (m, 1H); ¹³C NMR (CDCJyCD₂OD) 8 13.72, 22.40, 24.71, 28.84, 28.97, 29.08, 29.18, 29.41, 31.65, 34.1660.15, 60.99, 62.42, 63.17, 65.16, 65.30, 65.98, 73.24, 173,79; MS m/z 573 (M-H); IR (NaCl Neat) 3431, 1739, 1254, 1177 cm⁻¹

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Example 10 - Xenopus Oocyte Assay

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Xenopus cocytes which endogenously express PSP24 PLGFR were used to screen the newly designed and synthesized compounds for their LPA inhibitory activity.

Oocytes were obtained from xylazine-enesthetized adult Xenopus laevis frogs (Carolina Scientific, Burlington, NC) under aseptic conditions and prepared for experiment. Stage V-VI oocytes were denuded of the the follicular cell layer with type A collagenase treatment (Boehringer, IN) at 1.4 mg/ml in a Ca²⁺-free ovarian Ringers-2 solution ((OR-2) 82.5 mM NaCl, 2 mM KCl, 1 mM MgCl₃, 5mM HEPES, pH 7.5, with NaOH). Oocytes were kept in Barth's solution in an incubator between 17-20 °C and were used for 2-7 days after isolation.

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Electrophysiological recordings were carried out using a standard twoelectrode voltage-clamp amplifier holding the membrane potential at -60 mV (GeneClamp 500, Azon Instruments, CA). Test compounds were dissolved in MeOH, complexed with fatty acid free BSA, and diluted with frog Na⁺-Ringers solution (120 nM NaCl, 2 mM KCl, 1.8 mM CaCl₂, 5 mM HEPES; pH 7.0), which were applied through superfusion to the oocyte at a flow rate of 5 ml/min. Membrane currents were recorded with a NIC-310 digital oscilloscope (Nicolet, Madison, WI). Applications were made at intervals of 15 mins (minimum) to allow for the appropriate washout and recovery from desensitization.

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Figures 21-27 show the dose-dependent inhibition of LPA-induced chloride currents by compounds 56, 57, 66, and 92.

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Compound 36 was the best inhibitor among the non-phosphorylated derivatives. When compound 36 was injected intracellularly to see whether its

WO 01/71022

- 67 -

PCT/US01/08729

inhibitory effects were a result of its actions on the cell surface or whether the inhibition was a result of its actions within the cell, this intracellular application of 36 did not give any information as to its site of action. Hence, moving away from free hydroxy compounds (35-43), phosphorylated compounds (55-59) were synthesized to interact on the cell surface and to prevent the compounds from penetrating into the

Compounds 56, 57, 66, and 92 were inhibitors of LPA-induced chloride current in Xenopus oocyte. Compounds 56, 57, 66, and 92 were able to block the actions of LPA in a dose-dependent fashion. Moreover, washing the the Xenopus oocyte, there was a complete recovery of the LPA response; that experiment implies that compounds 56, 57, 66, 92 were able to inhibit the LPA-induced chloride currents in a reversible fashion. Compound 66 at 5 µM completely abolished the effect of LPA in Xenopus oocytes, with an IC₁₀ of about 1.2 µM (Figures 23 and 24). Moreover, when 66 was microinjected inside the cell (arrow, Figure 23B), followed by the extracellular application of LPA (10 nM), it failed to inhibit the LPA response; that extracellular nature.

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Compounds 35, and 37-43 were tested on Kenopus occytes, but the results were inconclusive. Compound 55 at 1 µM showed slight inhibition (38% against 2 nM LPA). In the SAP series, compounds 58 and 59 remain to be tested in the Kenopus occyte assay. In the bisphosphate series, compound 89 inhibited the LPA-induced response (59 % against 2 nM LPA). However, compounds 67 (threshold ~ 1 µM), 68 (threshold ~ 10 nM), and 85 (threshold ~ 10 nM) were able to elicit a response alone; compounds 86, 87, 88, 90, and 91 have yet to be evaluated.

Compound 56a was evaluated in the Kenopus occyte assay, 56a enhanced the LPA response when applied in combination with LPA. Compound 56a did not elicit a response at 2 µM (not shown), but at 10 µM, 56a was able to elicit a response on its own (Figure 26); that experiment suggests, that a free amino group is necessary for the inhibitory activity.

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Example 11 - HEY Ovarian Cells Migrations

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It is known that two LPA receptors, EDG-2 and EDG-7, are expressed in HEY ovarian cancer cells, so compounds \$6, \$6a, and 66 were evaluated for their ablity to inhibit LPA-induced cell motility (compound cone: 1µM against 0.1 µM LPA cone.).

- 68 -

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HEY ovarian cells were maintained in RPMI 1640 medium with 2 mM L-glutamine (GIBCO BRL) supplemented with 10% fetal bovine serum (FBS, Hyclone). All cells were synchronized to the Gy/G₁ stage by growing them to confluency for 2 days. The cells were replated and harvested for experiments when cells were about 50 -60% confluent on the flask. After removal of the cells from the flask, they were exposed for 5 min to 0.53 mM EDTA in PBS at 37°C. EDTA was neutralized with equal volume of RPMI 1640 plus 2 mM L-glutamine and 10% FBS. Cells were centrifuged at 800 rpm for 10 min at room temperature. Harvested cells were washed twice with RPMI 1640 with 2 mM L-glutamine medium and resuspended in the concentration of 1 x10° cells/ml, and then rested for 1 hr at 37°C.

A modified quantitative cell migration assay (Cat. # ECM500 from Chemicon, Temecula, CA) was used to test cell motility. The Chemicon chamber membrane was coated with fibronectin-containing pores of 8 microns in diameter. A 400 µl RPMI/2 mM L-glutamine containing either no inhibitors or inhibitors (1 µM) were pippetted into the lower chamber. About 5 x 10⁴ cells in RPMI 1640/2 mM L-glutamine were added to the top chamber. The 24-well plates with inserts were incubated for 4 hours in a 5% CO₂ incubator at 37°C. At the end of incubation, the chambers were removed to a fresh 24-well plate, and the cells on the inside chamber were removed by a swab several times and placed in the prepared Cell Stain Solution for 30 minutes at room temperature. At the end of incubation, Cell Stain Solution was removed from the wells. The chambers were washed 3 times with 1 mL PBS per well. After the final PBS wash, the chambers were examined to confirm proper cell morphology, and adherent cells were counted using an inverted microscope.

An effect of the newly synthesized compounds on the LPA-induced

migration of HEY ovarian cancer cell is shown in Figure 27. Compound 66 inhibited the LPA-induced cell motility by about 70%; however, compound 55 (marginally) and 56a potentiated the LPA-induced cell motility.

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Example 12 - Compound Cytotoxicity

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Im et al. (2000) and RT-PCR data showed the presence of PLGFR's in prostate cancer cell lines DU-145, PC-3, and LNCaP. Due to the promising inhibitory activity in *Xenopus* oocyte and the cell motility assay, the growth inhibitory effects of a number of compounds on DU-145, PC-3, and LNCaP prostate cancer cell lines were examined.

DU-145, PC-3, and LNCaP cells were propagated in 150 cm² flasks, containing RPMI-1640 or Dulbecco's modified Eagle media supplemented with 10%

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WO 01/71022

- 69 -

PCT/US01/08729

fetal bovine serum (FBS). Cells were removed from stock flasks using trypsin, centrifuged, resuspended in fresh media, and plated at a density of approximately 2,000 cells/well in 96-well culture plates. Final drug concentrations ranged from 0.05 to either 10 or 50 µM. Control experiments with no drug added (negative control) and 5-fluorouracil added (positive control) were performed in parallel. Media was removed and replaced at 48 hours to minimize the effects of drug degradation during the course of the experiment. After 96 hours drug exposure, cells were fixed by the addition of cold 50% trichlorosectic seid (TCA) and incubation at 4°C for 1 hour. Fixed cells were stained with sulforthodamine B (SRB), and cell number was determined by

comparison of absorbance at 540 nm, as compared to a standard curve of cell number versus absorbance. Experiments were performed in duplicate. Cell number as a percentage of control (untreated wells) was plotted versus drug concentration and the concentration that inhibited cell growth by 50% (IC₅₀) determined by nonlinear regression (WinNonlin, Pharsight Corporation).

Cytotoxicity studies performed on prostate cancer cell lines DU-145, PC-3, and LNCaP, together with the reference compounds 5F-uracil, LPA (18:1), SPH (13:0),

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LNCaP, together with the reference compounds 5F-uracil, LPA (18:1), SPH (13:0), SPP (13:0), and N-palmitoyl L-serine phosphoric acid (15:0), are shown in Table 3 below.

- 70 -

PCT/US01/08729

Table 3: Cytotoxicity of Synthesized Compounds on Prostate Cancer Cell Lines

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Compound	DU145	PC:3	LNCaP
Fluorouracil	6.8±3.3	10.2±4.1	2.8±1.6
LPA (18:1)	WA	28.5±6.3	WA
SPP (13:0)	> 10	WA	NA .
SPH (13:0)	13,9±1.1	11.7±2.3	5.7±2.1
N-palmitoyl-L-sorine (15:0)	WA	WA	WA
27	19.7±6.0	WA	10.9±2.7
38	38.9±8.9		٠
51	8.1±1.3	25.4±3.6	19.9±6.4
55	24.944.1	31.6±9.0	4.9±2.6
26	2.3±1.2	0.7±0.1	13.5±4.7
56a	0.7±0.1	WA	30.3±7.9
57	9.1±0.8	WA	10.7±2.1
99	NA	NA	3.1±3.2
29	WA	WA	25.2±12.3
89	WA	WA	29.3±21.7
88	Ą	NA	11.6±10.3
98	NA	NA	6
84	NA	NA	WA
88	NA	NA	4
68	WA	NA	7
96	> 50	WA	WA
16	42,2±1.9	WA	WA
92	WA	WA	WA

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*Cell number as a percentage of control (untreated wells) was plotted versus drug contentration and the concentration that inhibited cell growth by 50% (IC₅₀) determined by nonlinear regression (WinNonlin, Pharsight Corporation).

WA = Weak Activity; NA = No Activity;? = Maximum inhibition was 50%.

Compounds 55, 56, 56a, 66, and 85 exhibited a range of growth inhibitory activities. Compound 56 was a more potent inhibitor of DU-145 and PC-3 cell growth than 5-fluorouracil. Interestingly, 56a selectively inhibited DU-145 cell growth, but was less potent against PC-3 cells; compound 55 was a more potent inhibitor of LNCaP cell growth as against DU-145 and PC-3 cells. Compound 66 selectively inhibited LNCaP cell growth, but showed no activity on PC-3 and LNCaP cells. Compound 85 was the most active among the bisphosphates (sr-1 acyl).

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WO 01/71022

-71-

PCT/US01/08729

Discussion of Examples 1-12

Three seris of compounds were specifically synthesized and analyzed (35-43, 55-59, 66-68, and 85-92). The first and the second sets involve the amalganation of the endogenous inhibitors SPH and SPP with the synthetic inhibitor N-palmitoyl L-serine phosphoric acid, whereas the third series involves the bisphosphates. Compounds 56, 57, 66 and 92 were inhibitors of LPA-induced chloride currents in the *Xenopus* oocyte assay. Also, bisphosphates with shorter chain length at (3n-1) position were able to elicit chloride currents in *Xenopus* oocyte [67 (threshold ~ 1 µM), 68 (threshold ~ 10 nM), and 85 (threshold ~ 100 nM)]. Compound 66 was shown to inhibit the LPA-induced cell motility in HEY ovarian cancer cell lines. On evaluating the growth inhibitory effects of the above-synthesized compounds on DU-145, PC-3, and LNCaP prostate cancer cell lines, three highly potent and selective compounds (56, 56a, and 66) were discovered.

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The above data (Table 3) suggests that (i) compounds that contain an alcohol with no phosphate are less active (27 vs. 56), (ii) compounds with the protected phosphate moiety are less active (31 vs. 56), (iii) alkylation of the amine does not reduce activity (56a), (iv) the most potent bisphosphate has an ether linkage at the *sn*-1 position, (v) decreasing the chain length in the SAP series (55 vs. 56) decreased the potency towards DU-145 and PC-3 (however, it was more potent against LNCaP cells), (vi) on decreasing the chain length for the bisphosphate (*sn*-1 alkyl) compounds, potency decreased, though selectivity towards LNCaP cell remained, and (vii) substitution at *sn*-1 position (azyl vs alkyl) did not increase the potency. The target site for these molecules is likely on the cell membrane (e.g., a membrane-spanning receptor), because the polar phosphate derivatives are unlikely to easily cross the cell membrane (although there exists the possibility that an active transport system could exist). These results suggest that differences in PLGFR's or downstream signal transduction events may play a significant role in the growth inhibitory properties of these compounds in prostate cancer cells.

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Example 13 – Preparation and Characterization of Stable Cell Lines Expressing Edg. 2, Edg. 4, and Edg. 7

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In an effort to develop selective aniagonists to the Edg-2, -4, and -7 receptors, a system for screening potential compounds was first established. RH7777 cells were chosen as a model system since they have been reported to be non-responsive to LPA in a variety of cellular assays and were found to be devoid of mRNA for any of the known Edg receptors (Fukushima et al., 1998). Stable cells lines

-72-

PCT/US01/08729

transfected with the EDG receptors, as well as control cell lines transfected with empty vector, were established in RH7777 cells.

The resulting clones were screened by monitoring intracellular Ca²⁺ transients, and by RT-PCR. This screening process led to the identification of at least three positive cell lines expressing Edg-2 and -7, while no positive cell lines expressing Edg-4 could be identified. Vector transfected cells were also found to be non-responsive to LPA. Although stable clones expressing Edg-4 were not isolated, the transient expression of Edg-4 resulted in the LPA-mediated activation of intracellular Ca²⁺ transients, demonstrating that the construct was functionally active in these cells. The stable Edg-4 cell line used in these experiments was isolated and characterized by Im et al., who kindly provided us with the same clone (Im et al., 2000).

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The cell lines were further characterized in an effort to identify a suitable assay for screening potential antagonists. LPA-ellicited activation of ERK 1/2 was seen in Edg-2 and transient Edg-4 expressing cells, whereas ERK 1/2 was not activated in Edg-7 expressing cells. LPA elicited Ca²⁺ transients in all stable cell lines expressing Edg-2, -4, and -7. Dose response curves revealed EC₅₀ values of 378 ± 53, 998 ± 67, and 214 ± 26 nM for Edg-2, -4, -7 expressing cells, respectively (Figures 28A-C). Because the EC₅₀ value determined in the stable Edg-4 clone was different from that previously reported, a dose response curve was also established for cells transiently expressing Edg-4 (Figure 28B, An et al., 1998a; An et al., 1998b), which yielded an EC₅₀ value of 186 ± 39.

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The ability of LPA to stimulate DNA synthesis in the stable cell lines was examined by measuring the incorporation of ³H-thymidine. Neither wild type, nor the vector transfected RH7777 cells showed an increase in ³H-thymidine incorporation following a 24 hr incubation with 10 μM LPA, which is in contrast to a previous report that LPA is mitogenic in these cells. Edg-2 expressing cells showed a 1.8-fold increase in ³H-thymidine incorporation, whereas Edg-4 and –7 expressing cells did not show an increase in ³H-thymidine incorporation, as compared to control cells.

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Example 14 – Short Chain Phosphatidates Activity on Edg-2 and Edg-7 Receptors

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Since Ca²⁺ transients were elicited in all three stable cell lines expressing Edg-2, -4, and -7 (Figures 28A-C), this assay was used for screening potential antagonists. In an effort to identify selective antagonists for the LPA activated members of the Edg receptor family, Edg-2, -4, and -7, the structural features of the LPA pharmacophore were relied upon as a starting point. Short-chain (8:0)

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WO 01/71022

- 73 -

PCT/US01/18729

LPA or a mixture of LPA (8:0) and LPA (18:1) were tested as inhibitors of Edg-2, -4, or -7. When the cells were challenged with the mixture of LPA 8:0 and LPA 18:1, Ca²⁺ responses were not effected in any of the three stable cell lines (see Figures 30A-C, 31A-C, and 32A-B). LPA 8:0, alone, was unable to elicit Ca²⁺ responses in any of the cells, at concentrations as high as 10 μM.

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Based on these results, applicants hypothesized that a modification of the LPA pharmacophore, which sterically restricted the mobility of the fatty acid chain, might also effect its ligand properties. For this reason, we tested compounds with a second short-chain fatty acid at the sn-2 position were also tested. Such short-chain phosphatidates have increased hydrophobicity over the corresponding short-chain LPA, which could exert constraints on their interaction with the ligand-binding pocket of the receptor.

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in Edg-4 expressing cells were unaffected by DGPP 8:0 (Figure 29B). Because of the Similar observations were obtained with PA 8:0 in each of the assays described above with Edg-4. Consistent with results from experiments in stable cells, Ca2+ responses expressing cells were completely abolished (Figure 29C). In contrast, Ca²⁺ responses inhibited by approximately 50% (Figure 29A), whereas the responses in Edg-7 LPA in the stable cell lines. The Ca2+ responses in Edg-2 expressing cells were DGPP were prepared and tested as an inhibitor of Edg-2, -4, or -7. Figures 29A-D an agonist of the Edg receptors (see below). With this similarity in mind, short-chain naturally occurring lipids which share some key chemical properties with the LPA for DGPP 8:0 (see below). were not effected by DGPP 8:0 in cells transiently expressing Edg-4 (Figure 29D). show the effect of a 10-fold excess of DGPP (8:0) on the Ca²⁺ responses elicited by pharmacophore, having an ionic phosphate group(s) and fatty acid chains. Neither is (Figure 29B), DGPP 8:0 was similarly tested on cells that were transiently transfected discrepancy in EC30 values for the stable and transient expression of Edg-4 Phosphatidic acid (PA) and diacylglycerol pyrophosphate (DGPP) are

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Inhibition curves were determined in cells expressing Edg-2 and -7, using increasing concentrations of DGPP 8:0, while the concentration of LPA was kept constant at the EC₅₀ relative to the receptor studied. IC₅₀ values of 285 ± 28 nM for Edg-7 (Figure 30A) and 11.0 ± 0.68 µM for Edg-2 (Figure 31A) were determined from the curves. Using a constant amount of DGPP 8:0 near to the IC₅₀ value (250 nM for Edg-7, 3 µM for Edg-2), the dose response curves for both Edg-7 (Figure 30B) and Edg-2 (Figure 31B) were shifted to the right, indicating a competitive mechanism of inhibition.

PCT/US01/IB729

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LM, all decreased the responses to 50, 19, and 64% of control, respectively. When the Ca22 response, in agreement with the results from the stable cell line. In contrast to the response at concentrations up to 10 µM, in cells stably or transiently expressing Edg-4 (Figure 31C). Octyl chain length analogs of DGPP, PA, and DAG, when used at 10 control, respectively. To confirm the results obtained from the stable clone, the lipid Edg-4, none of the short- or long-chain lipids had an inhibitory effect, whereas both the short-, nor the long-chain species of DGPP or PA had an inhibitory effect on the panel was tested on cells transiently expressing Edg-4 (Figure 32B). Again, neither (Figures 32A-B). When these lipids were assayed in the stable cell line expressing transient expression of Edg 4. Neither species of PA when applied alone, elicited a effect, whereas PA 18:1 maintained a modest inhibitory effect, decreasing the ${
m Ca}^{2+}$ chain length was increased to 18:1, DGPP and DAG no longer had an inhibitory PA 8:0 and 18:1 significantly increased the Ca2+ responses, to 162 and 137% of response by 18%. The panel of lipids was also tested on Edg-4 expressing cells stable Edg-4 clone, neither PA analog enhanced the Ca2+ response in cells with The same set of lipids was tested on Edg-2 expressing cells

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The effect of DGPP 8:0 on cells that endogenously express LPA receptors was also examined. DGPP 8:0 was found to inhibit the Ca²⁺ mediated, inward Cl' currents elicited by LPA in *Xenopus* cocytes with an IC;0 of 96 ± 21 nM (Figure 33A). In the presence of a 200 nM concentration of DGPP 8:0, the dose response curve for LPA 18:1 was shifted to the right, indicating a competitive mechanism of action as found in Edg-2 and -7 clones (Figure 33B). To examine whether DGPP 8:0 acts through an intracellular or extracellular mechanism, DGPP 8:0 was injected intracellularly and the occyte was exposed to LPA 18:1. Figure 32C shows that following the intracellular injection of DGPP 8:0, estimated to reach a concentration > 300 nM, the extracellular application of 5 nM LPA 18:1 elicited a

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WO 01/71022

- 75 - PCT/US01/08729

response equal in size to that of the control. In comparison, the response normally elicited by LPA 18:1 was completely inhibited when DGPP 8:0 was applied extracellularly (Figure 33C). The inhibitory effect of DGPP 8:0 was reversible, as after a 10-min washing the response recovered to control level (Figure 33C).

To show the specificity of DGPP 8:0 for the LPA receptors expressed in the oocyte, the expression of neurotransmitter receptors was induced by the injection of polyA+ mRNA from rat brain. This resulted in the expression of the G-protein coupled receptors for serotonin and acetycholine, which are not expressed in non-injected oocytes. These neurotransmitters activate the same inositol trisphophate. Ca^{2*} signaling pathway that is activated by LPA (Tigyi et al., 1990). In these oocytes, DGPP 8:0 did not inhibit either serotonin- or earbachol-elicited responses, demonstrating the specificity of DGPP 8:0 for the LPA receptors. PA 8:0 when used at similar concentrations was also effective at inhibiting the LPA-elicited responses in the oocytes.

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The effect of DGPP 8:0 on LPA-elicited responses was also examined in mammalian systems that endogenously express LPA receptors. NHHJT3 cells were screened by RT-PCR for the presence of mRNA for the Edg and PSP24 receptors. Figure 34A shows that in NHJT3 cells mRNA transcripts for Edg-. -5, and PSP24 were detected. To show that DGPP 8:0 was specific in inhibiting LPA-elicited but not SIP-elicited Ca²⁺ responses, NHJT3 cells were exposed to 100 nM LPA or SIP in the presence of 10 µM DGPP 8:0. As shown in Figure 34B, DGPP 8:0 significantly inhibited the LPA-elicited Ca²⁺ responses, whereas the SIP-elicited response was not effected.

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LPA has been shown to be generated from and play a role in ovarian cancer (Xu et al., 1995a). Therefore, DGPP 8:0 was also tested on HEY ovarian cancer cells to determine if it had an effect on a therapeutically relevant target. Figure 34D shows that DGPP 8:0 inhibited the LPA-elicited Ca²⁺ response to 12% of control, whereas DGPP 18:1 had no effect. Likewise, PA 8:0 inhibited the Ca²⁺ response to 6% of control, whereas PA 18:1 had no effect. HEY express mRNA transcripts for Edg-1, -2, -5, -7 receptors (Figure 34C).

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Example 15 - Inhibition of NIH3T3 Cell Proliferation

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The hallmark effect of a growth factor is its ability to elicit cell proliferation. Since LPA has been shown to stimulate the proliferation of a variety of different cell types (Goerzl et al., 2000), the ability of DGPP 8:0 to inhibit cell proliferation was examined in NH3T3 cells. Figure 35 shows that DGPP 8:0

- 76 -

PCT/US01/08729

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significantly inhibited the LPA-induced proliferation of NIH3T3 cells, reducing cell number to control levels, whereas it had no effect on the solvent-treated control cells. To define the structure-activity relationship for the inhibitory effect of DGPP 8:0, the short- and long-chain species of DGPP, PA, and DAG were included in the assay. As shown in Figure 35, none of the lipids included in the test panel had a significant inhibitory or stimulatory effect on the solvent-treated control cells. Only DGPP 8:0 inhibited the LPA-induced proliferation. Neither DGPP 18:1, nor long- and short-chain PA and DAG had an effect on the LPA-induced proliferation. Interestingly, PA 8:0 had no significant inhibition in this assay.

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Discussion of Examples 13-15

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RH7777 cells were used for heterologous expression of Edg-2, -4, and -7 receptors to screen potential antagonists. Based on our previous computational modeling of the Edg receptors (Parrill et al. 2000) and the available structure-activity data (Jalink et al., 1995), the above experimental results demonstrate that the short-chain phosphatidate DGPP 8:0 is a selective, competitive antagonist of Edg-7, with an IC₁₀ value of 285 ± 28 nM. The same molecule was found to be a poor inhibitor of Edg-2, with an IC₂₀ value of 11.0 ± 0.68 μM, whereas it did not inhibit Edg-4. DGPP 8:0 inhibited the endogenous LPA response in Xenopus occytes with an IC₂₀ value of 96 ± 21 nM. PA 8:0 showed similar inhibitory properties. Therefore, these short-chain phosphatidates show a 40-100-fold selectivity for Edg-7 over Edg-2.

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The above results with short-chain phosphatidates confirm those of Bandoh et al. (2000) who demonstrated that LPA, with an acyl chain-length of twelve carbons or less, does not elicit responses in insect cells expressing Edg-2, -4, or -7. As demonstrated above, LPA 8:0 was neither an agonist nor an antagonist of Edg-2, -4, or -7 in a mammalian expression system. Edg-7 has a 10-fold preference for LPA with the faity acid chain esterified to the sr-2, versus the sr-1 position (Bandoh et al., 2000). Therefore, the distance of the hydrocarbon chain relative to the phosphate moiety, does not abolish the binding to and activation of the receptor. Edg-7 also shows a preference for long-chain, unsaturated fatty acids over their saturated counterparts. The presence of an ether linkage or vinyl-ether side chain also decreased the EC₂₀ by two orders of magnitude (Bandoh et al., 2000). Moreover, there is an optimal hydrocarbon chain-length of 18 carbons, whereas 20 carbon analogs were weaker agonists. These pharmacological properties of Edg-7 suggest that receptor activation is dependent upon the chain length, as well as the flexibility of the side chain (ester vs. ether linkage).

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WO #1/71022 - 77 -

PCT/US01/18729

moiety, which does not change the negatively charged character of the anchoring members of the Edg family. The second residue, arginine 292, occurs at a position not to be able to activate the receptors due to their truncated hydrocarbon chains. The Because of the relative tolerance of the sn-1 and sn-2 substitution of the fatty acids by anchor and the hydrophobic tail, are required for agonist activation. In support of this not sufficient for ligand binding and activation (Parrill et al., 2000). It was Moreover, the interaction between the receptor and the hydrocarbon chain, itself, was predicted to interact with the hydroxyl moiety of LPA. Alanine replacement of this a glutamine at the corresponding site in Edg-2, -4, and -7. This glutamine residue is residue, glutamate 121, is not conserved amongst the LPA-specific Edg receptors, with region, but rather increases the charge. structural mobility of the acyl chains in the phosphatidates is also limited by the designated the hydrophobic tail as the "switch" region of the PLGF pharmacophore. activate Edg-2, -4, or -7, underlying the importance of the interaction between the hypothesis, the above results demonstrate that the short-chain LPA 8:0 was not able to hypothesized, therefore, that a combination of interactions, involving both the ionic and these three residues is necessary for ligand binding in Edg-1 (Parrill et al., 2000). that the ionic interaction between the charged moieties of the PLGF pharmacophore residue has led to a loss of ligand binding and activation of the receptor, suggesting where all Edg family members except Edg-8 have a nearby cationic residue. The third charged residues that are required for ligand binding. One of these residues, arginine adjacent fatty acid moiety. Applicants also explored the effects of a pyrophosphate these receptors, applicants focused on short-chain phosphatidates which were believed hydrophobic tail and the ligand binding pocket. As a result, applicants have 120, which is predicted to interact with the phosphate group, is conserved in all of the Computational modeling of the Edg-1 receptor has identified three

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This conceptual drug design was tested on clonal cell lines expressing the Edg-2, -4, and -7 receptors. The pharmacological properties of DGPP 8:0 and PA 8:0 were found to be dramatically different between the three receptors. Both molecules were effective at inhibiting Edg-7, whereas they were more than an order of magnitude less effective on Edg-2. Neither molecule was effective on Edg-4. DGPP 8:0 was also found to be a competitive inhibitor of both Edg-2 and -7, displacing the dose response curves to the right with a subsequent increase in the EC₅₀ values for LPA on both receptors. The lack of agonist activity of the corresponding long-chain species of PA and DGPP, highlights the constraints that prevail in the binding pocket, is

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- 78 -

supported by the lack of inhibition by DAG 8:0, although its cellular effects are likely confounded by its intracellular actions on other molecular targets, such as PKC.

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Both PA and DGPP are naturally occurring phospholipids. DGPP (8:0) was discovered in 1993 as a novel lipid in plants and is a product of the phosphorylation of PA by phosphatidate kinase (Wissing and Behrbohm, 1993; Murnik et al., 1996). DGPP has been identified in bacteria, yeast and plants, but not in mammalian cells. Recent studies have shown that DGPP activates macrophages and stimulates prostaglandin production through the activation of cytosolic phospholipase A₃, suggesting a role for DGPP in the inflammatory response (Balboa et al., 1999; Balsinde et al., 2000). These authors ruled out the possibility that these effects were mediated through LPA receptors. The above results with the long-chain DGPP and PA analogs confirmed this notion, as these compounds did not possess agonist properties in the Edg receptor expressing cell lines at concentrations up to 10

The effect of short chain phosphatidates was also examined on LPA receptors expressed endogenously in three different cell types. DGPP 8:0 and PA 8:0 were found to be effective inhibitors of LPA-elicited Cl currents in *Xenopus* oocytes. In order to determine the site of action, DGPP 8:0 was injected into oocytes followed by an extracellular application of LPA. DGPP 8:0 was only effective at inhibiting the LPA-elicited Cl currents when applied extracellularly, demonstrating that it exerts its amagonist effect on the cell surface. The specificity of DGPP 8:0 for LPA receptors was demonstrated in oocytes and NIH3T3 cells. In these cells, DGPP 8:0 was only effective at inhibiting the LPA-elicited Ca^{2*} responses and not the responses elicited by S1P, acctycholine, or serotonin.

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RT-PCR analysis revealed that only Edg-2, and not Edg-4, or -7 is expressed in NIH3T3 cells. In NIH3T3 cells, DGPP 8:0, at a high 100-fold excess, only inhibited the Ca²⁺ responses by 40%. This degree of inhibition parallels that seen in the stable cell line expressing Edg-2, where it was also a weak inhibitor. When short-chain DGPP and PA were evaluated on HEY ovarian cancer cells, at a 10-fold excess over LPA, both were effective inhibitors, whereas neither long-chain molecule had any effect. RT-PCR revealed that the predominant mRNA was for Edg-7 in HEY cells, whereas only a trace of Edg-2 mRNA was detected. This degree of inhibition parallels that seen in the stable cell line expressing Edg-7, where both DGPP 8:0 and PA 8:0 were effective inhibitors.

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Both short chain phosphatidates were evaluated for their ability to block the LPA-induced proliferation of NIH3T3 cells. DGPP 8:0 effectively inhibited the LPA-induced proliferation, while the long-chain DGPP did not. Although PA 8:0

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WO 01/71022

- 79 -

was effective at inhibiting the Ca* responses, it was not effective at inhibiting cell proliferation. These results are in agreement with a previous report that PA (12:0) did not inhibit the mitogenic effect of PA 18:1 (van Corven et al., 1992). The stability of the molecules in long-term assays is a concern, since lipid phosphatases might inactivate the antagonist. The fact that both PA and DAG failed to inhibit the proliferation suggests that DGPP 8:0 is likely to be more stable for the duration of this assay. The stability of DGPP has also been demonstrated by Balboa et al. (1999), who reported that DGPP was not metabolized during the course of their experiments.

DGPP 8:0 provides an important new tool for the field in studying, not only the Edg receptors but also other PLGF receptors. The concept of an ionic anchor and hydrophobic switch of the PLGF pharmacophore derived from computational modeling of the Edg family should assist the design and synthesis of new inhibitors.

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Example 16 - Synthesis of Straight-Chain Phosphate Intermediates 101-105

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Compound 101: Phosphoric acid dibenzyl ester butyl ester

74 mg (1.00 mmol) of anhydrous n-butanol and 365 mg (5.17 mmol) of 1H-tetrazole were dissolved in 34 mL of anhydrous methylene chloride in a 100 mL round-bottom flask. A solution of 0.895 g (2.58 mmol) of diberazyl-N/M-diisopropyl phosphoramidite in 5 mL of anhydrous methylene chloride was added via a syringe under an argon atmosphere with stirring. The reaction mixture was stirred at room temperature for 2 hrs. The reaction mixture was then cooled in a isopropyl alcohol/dry ice bath at ~38 °C. 0.815 g (3.43 mmol) of 32 % peracetic acid in 28 mL of anhydrous methylene chloride were added dropwise via an addition fumel. After the addition, the temperature of the reaction mixture was raised to ~0 °C with an ice bath. The reaction mixture was strated in the ice bath for 1 hr. The reaction mixture was transferred to a separatory furnel and diluted with 200 mL of methylene chloride. The organic layer was washed with 10% sodium metabisulfite (2 x 40 mL), saturated sodium bicarbonate (2 x 40 mL), water (30 mL), and brine (40 mL). The organic layer

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amount of impurity from excess phosphorylating reagent) as a clear oil. ¹H NMR (CDCi₃) 0.88 (t, J = 7.2 Hz, 3H, CH₃), 1.34 (seatet, J = 7.2 Hz, 2H, OCH₃CH₃CH₃CH₃), 1.59 (quintet, J = 6.6 Hz, 2H, OCH₃CH₃CH₃CH₃), 3.99 (dt, J = 6.6 Hz, 2H, OCH₃CH₃CH₃CH₃CH₃), 5.02 (d, J = 1.8 Hz, 2H, OCH₃Ah), 5.03 (d, J = 2.1 Hz, 2H, OCH₃Ah), 7.35 (br s, 10H, 2 x ArH); ¹³C NMR (CDCl₃) 13.55, 18.60,

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dryness. The crude product was then purified by silica gel chromatography using 1:1

nexanes/ethyl acetate as the eluent to afford 101 (309 mg which contained a slight

was dried with anhydrous sodium sulfate, filtered, and concentrated under vacuum to

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PCT/US01/08729

mode): [M + 23Na] at m/x 357.3. 32.16 (d, $J_{C,P} = 6.8$ Hz), 67.72 (d, $J_{C,P} = 6.1$ Hz), 69.13 (d, $J_{C,P} = 5.5$ Hz), 127.90, 128.47, 128.55, 136.00 (d, J_{C,P}= 6.8 Hz); ³¹P NMR (CDCl₃) 16.84; MS (positive

Compound 102: Phosphoric acid dibenzyl ester octyl ester

OCH2CH2(CH2)5CH3), 1.60 (quintet, J= 6.9 Hz, 2H, OCH2CH2(CH2)5CH3), 3.98 (dt, J (CDCl₃) 16.83; MS (positive mode): $[M + {}^{23}Na]^{+}$ at m/z 413.4. 69.12 (d, $J_{C,P}$ = 5.5 Hz), 127.90, 128.47, 128.56, 135.97 (d, $J_{C,P}$ = 6.9 Hz); ³¹P NMR 22.62, 25.38, 29.06, 29.14, 30.17 (d, $J_{CP} = 6.9 \text{ Hz}$), 31.75, 68.05 (d, $J_{CP} = 6.2 \text{ Hz}$), (d, J = 2.4 Hz, $2H OCH_2Ar$), 7.34 (br s, 10H, $2 \times ArH$); ^{13}C NMR (CDCl₃) 14.09, = 6.6 Hz, 6.9 Hz, 2H, OCH₂CH₂(CH₂)₅CH₃), 5.02 (d, I = 2.1 Hz, 2H, OCH₂Ar), 5.05 90%) as a clear oil. ¹H NMR (CDCl₃) 0.88 (t, J=6.9 Hz, 3H,CH₃), 1.24 (br s, 10H, chromatography using 7:3 hexanes/ethyl acetate as the eluent to afford 102 (351 mg, analogous to that for 101 was performed. The crude product was purified by silica gel 130 mg (1.00 mmol) of anhydrous n-octanol were used and a procedure

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Compound 103: Phosphoric acid dibenzyl ester dodecyl ester

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30.18 (d, $J_{CP} = 7.0$ Hz), 31.92, 68.05 (d, $J_{CP} = 6.1$ Hz), 69.12 (d, $J_{CP} = 5.4$ Hz), ArH); 13C NMR (CDCl3) 14.13, 22.69, 25.38, 29.12, 29.35, 29.49, 29.56, 29.63, (d, J = 2.1 Hz, 2H, OCH₂Ar), 5.05 (d, J = 2.1 Hz, 2H, OCH₂Ar), 7.34 (br s, 10H, 2×10^{-1} $OCH_2CH_2(CH_2)_2CH_3$, 3.98 (td, J = 6.9 Hz, 6.6 Hz, 2H, $OCH_2CH_2(CH_2)_2CH_3$), 5.02 (br s, 18 H, OCH₂CH₂(CH₂)₉CH₃), 1.60 (quinter, J = 6.9 Hz, 2H, (361 mg, 81%) as a clear oil. 'H NMR (CDCl₃) 0.88 (t, J = 7.2 Hz, 3H, CH₃), 1.24 silica gel chromatography using 7:3 hexanes/ethyl acetate as the eluent to afford 103 procedure analogous to that for 101 was utilized. The crude product was purified by 186 mg (1.00 mmol) of anhydrous n-butanol were employed and a

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Compound 104: Phosphoric acid dibenbyl ester octadecyl ester

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(positive mode): [M + 23Na] at m/z 469.1.

127.89, 128.46, 128.55, 135.97 (d, J_{C,P}=6.8 Hz); ³¹P NIMR (CDCl₃) 16.84; MS

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(d, J = 2.1 Hz, 2H, OCH₂Ar), 5.05 (d, J = 2.1 Hz, 2H, OCH₂Ar), 7.34 (br s, 10H, 2 x Hz, 3H, CH₃), 1.25 (br s, 30H, OCH₂CH₂(CH₂)₁₅CH₃), 1.60 (quintet, J = 6.9 Hz, 2H, 89%) as a hygroscopic white solid: mp 32-33 °C; ¹H NMR (CDCl₃) 0.88 (t, J = 6.9 OCH2CH2(CH2)15CH3), 3.98 (td, J= 6.6 Hz, 6.9 Hz, 2H, OCH2CH2(CH2)15CH3), 5.02 chromatography using 7:3 hexanes/ethyl acetate as the eluent to afford 104 (474 mg. as for 101 was employed. The crude product was purified by silica gel 270 mg (1.00 mmol) of octadecanol were used and the same procedure

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WO 01/71022 <u>.</u>

PCT/US01/I88729

29.72, 30.20 (d, $J_{CP} = 6.9 \text{ Hz}$), 31.94, 68.06 (d, $J_{CP} = 6.1 \text{ Hz}$), 69.14 (d, $J_{CP} = 5.4 \text{ Hz}$). (positive mode): $[M + {}^{23}N_B]^{\dagger}$ at m/z 553.3. 127.90, 128.47, 128.55, 136.00 (d, J_{C,P}=6.8 Hz); ³¹P NMR (CDCl₃) 16.83; MS ArH); ¹³C NMR (CDCl₃) 14.12, 22.70, 25.40, 29.13, 29.38, 29.51, 29.58, 29.68,

Compound 105: Phosphoric acid dibenzyl ester docosanyl ester

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29.71, 30.18(d, $J_{CP} = 6.9 \text{ Hz}$), 31.93, 68.06 (d, $J_{CP} = 6.0 \text{ Hz}$), 69.13 (d, $J_{CP} = 5.6 \text{ Hz}$) 2 x ArH); ¹³C NMR (CDCl₃) 14.13, 22.70, 25.39, 29.12, 29.37, 29.50, 29.57, 29.66 6.9 Hz, 3H, C出), 1.25 (br s, 38H, OCH2CH2(CH2)19CH3), 1.60 (quintet, J=6.9 Hz, 88%) as a hygroscopic white solid: mp 43.5-44.5 °C; ¹H NMR (CDCl₃) 0.88 (t, J= chromatography using 7:3 hoxanes/ethyl acetate as the elueut to afford 105 (516 mg. (positive mode): [M + 23Na] at m/z 609.3. 5.02 (d, J = 2.4 Hz, 2H, OCH_2Ar), 5.05 (d, J = 2.4 Hz, 2H, OCH_2Ar), 7.35 (br s, 10H) 2H, $OCH_2CH_2(CH_3)_{19}CH_3$), 3.98 (td, J = 6.6 Hz, 6.6 Hz, 2H, $OCH_2CH_2(CH_3)_{19}CH_3$), procedure to that for 101 was used. The crude product was purified by silica gel 127.89, 128.47, 128.55, 135.98 (d, $J_{CP} = 6.9 \text{ Hz}$); ³¹P NMR (CDCl₃) 16.83; MS 327 mg (1.00 mmol) of docosanol were employed and an analogous

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Example 17 -Synthesis of Straight-Chain Phosphate Compounds 106-110

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Compound 106: Phosphoric acid monobutyl ester

Hz, 6.6 Hz, 2H, OCH2CH2CH2CH3); ¹³C NMR (CDCI3/ MeOH-da) 13.71, 19.02, OCH2CH2CH3CH3), 1.66 (quintet, J = 6.9, 2H, OCH2CH2CH3CH3), 3.99 (td, J = 6.6 evaporated under vacuum leaving behind 70 mg (86%) of a yellow oil 106. ¹H NMR vacuum through a pad of celite which was washed with methanol. The solvent was vessel at room temperature for 8 hrs. The reaction mixture was then filtered by apparatus and a hydrogen atmosphere of ~ 50 psi was maintained inside the reaction MS (negative mode): [M - 1]" at m/z 153.0. 32.72 (d, $J_{C,P} = 7.2 \text{ Hz}$), 66.86 (d, $J_{C,P} = 5.5 \text{ Hz}$); ³¹P NMR (CDClyMeOH-d₄) 18.84; (CDCl₃/MeOH-d₄) 0.95 (t, J = 7.2 Hz, 3H, CH₃), 1.43 (sextet, J = 7.5 Hz, 2H, 200 mg of 10% Pd/C was added. The vessel was connected to a hydrogenation methanol in a thick-walled pressure vessel. The vessel was purged with argon and \sim 200 mg (0.60 mmol) of 101 were dissolved in 30 mL of anhydrous

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Compound 107: Phosphoric acid monooctyl ester

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analogous to that for 106, 100 mg (93%) of a white/yellow tacky solid 107 was 200 mg (0.51 mmol) of 102 were employed and using a procedure

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PCT/US01/08729

22.98, 25.89, 29.57, 29.58, 30.76 (d, $J_{CP} = 7.3 \text{ Hz}$), 32.18, 67.16 (d, $J_{CP} = 5.2 \text{ Hz}$); ³¹P solated. ¹H NMR (CDCl₃/MeOH-4₄) $0.89 (t, J = 6.9 \text{ Hz}, 3H, C\underline{H}_3)$, 1.29 (br s, 10H, OCH1CH2(CH2)5CH3), 1.67 (quintet, J = 6.9 Hz, 2H, OCH2CH2(CH2)5CH3), 3.97 (dt, 1=6.6 Hz, 6.6 Hz, 2H, OCH2CH2(CH2); 13C NMR (CDCIs/McOH-42) 14.18, NMR (CDCl₃/MeOH-4₄) 20.55; MS (negative mode): [M - 1]^{*} at m/z 209.1.

Compound 108: Phosphoric acid monododecyl ester

OCH2CH2(CH2), CH3), 1.67 (quintet, J = 6.6 Hz, 2H, OCH2CH2), CH2), 3.97 (dt, J 22.98, 25.84, 29.57, 29.67, 29.89, 29.92, 29.96, 29.98, 30.69 (d, J_{Cr} = 7.4 Hz), 32.25, 200 mg (0.45 mmol) of 103 were employed and a procedure the same = 6.6 Hz, 6.6 Hz, 2H, OCH2CH2(CH3), CH3); 13C NMR (CDC1, MeOH-d4) 14.21, 67.22 (d, J_{GP}= 5.7 Hz); ³¹P NMR (CDCl₂/MeOH-d₄) 21.22; MS (negative mode): as that for 106 was used to afford 112 mg (94%) of a white solid 108. 'H NMR (CDCls/McOH-ds) 0.88 (t, J = 6.6 Hz, 3H, CH3), 1.27 (br s, 18 H, [M - 1] at m/z 265.0.

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Compound 109: Phosphoric acid monooctadecyl ester

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hat of 106 was employed which yielded 104 mg (79%) of a white solid 109. HNMR 14.26, 23.14, 26.01, 29.74, 29.84, 30.06, 30.09, 30.16, 30.87 (d, J_{CP} = 7.2 Hz), 32.42, 200mg (0.38 mmol) of 104 were used and an analogous procedure to 57.32 (d, J_{CP} = 5.8 Hz); ³¹P NMR (CDClyMeOH-d₄) 21.69; MS (negative mode) OCH₂CH₂(CH₂)₁₅CH₃), 1.68 (quintet, J = 6.9 Hz, 2H, OCH₂CH₂(CH₂)₁₅CH₃); 3.98 (dt, J = 6.6 Hz, 6.9 Hz, 2H, OCH2CH2(CH2)15CH3); 13C NMR (CDC13/MeOH-d4) (CDCl₃/MeOH-4,) 0.89 (t, J = 6.9 Hz, 3H, CH₃), 1.27 (br s, 30H, M - 1J at m/z 349.1.

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Compound 110: Phosphoric acid monodocosyl ester

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200 mg (0.34 mmol) of 105 were employed and the same procedure as 32.29, 67.27 (d, J_{CP} = 5.6 Hz); ³¹P NMR (CDCl₂/MeOH-d₄) 20.66; MS (negative OCH2CH2(CH2)19CH3), 1.66 (quintet, J = 6.9 Hz, 2H, OCH2CH2(CH2)19CH3), 3.97 (4, 1 = 6.6 Hz, 6.6 Hz, 2H, OCH2CH2(CH2)), CH3); 13C NMR (CDCI3/MeOH-4,) 14.22, 23.01, 25.87, 29.61, 29.71, 29.93, 29.97, 30.04, 30.73 (d, J_{C,P} = 7.4 Hz), hat for 106 was used yielding 98 mg (71%) of a white solid 110. ¹H NMR (CDCl₃/MeOH-d₄) 0.88(t, J = 6.9 Hz, 3H), 1.26 (br s, 38H, node): [M - 1] at m/z 405.1.

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WO 01/71022

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PCT/US01/08729

Straight-Chain Phosphate Compounds 106-110 Example 18 -

occytes were denuded of the the follicular cell layer with type A collagenase treatment (Boehringer, IN) at 1.4 mg/ml in a Ca2+-free ovarian Ringers-2 solution ((OR-2) 82.5 were kept in Barth's solution in an incubator between 17-20 °C and were used for 2-7 obtained from xylazine-anesthetized adult Xenopus laevis frogs (Carolina Scientific, used to screen compounds 106-110 for their LPA inhibitory activity. Oocytes were Burlington, NC) under aseptic conditions and prepared for experiment. Stage V-VI mM NaCl, 2 mM KCl, 1 mM MgCl2, 5mM HEPES, pH 7.5, with NaOH). Oocytes Xenopus oocytes which endogenously express PSP24 PLGFR were lays after isolation.

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were made at intervals of 15 mins (minimum) to allow for the appropriate washout and (GeneClamp 500, Axon Instruments, CA). Test compounds were dissolved in MeOH, through superfusion to the oocyte at a flow rate of 5 ml/min. Membrane currents were complexed with fatty acid free BSA, and diluted with frog Na +-Ringers solution (120 Electrophysiological recordings were carried out using a standard tworecorded with a NIC-310 digital oscilloscope (Nicolet, Madison, WI). Applications aM NaCl, 2 mM KCl, 1.8 mM CaCl2, 5 mM HEPES; pH 7.0), which were applied electrode voltage-clamp amplifier holding the membrane potential at -60 mV ecovery from desensitization.

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ibout 10.2 nM. Figure 37 compares the EC3 values for positive control solution (LPA lone), 25 nm, and a solution containing LPA and 100 nM of compound 108, 343 nM. rurrents, although compound 107 displayed a similar efficacy with an ${\rm IC}_{30}$ value of chain alkyl groups showed decreasing efficacy in inhibiting LPA-induced chloride naving an IC30 value of about 8.1 nM. Compounds with shorter or longer straightchloride currents by compounds 106-110. Compound 108 was the best inhibitor, Thus, compound 108 effectively inhibits LPA signalling of PSP24 receptors in Figure 36 shows the dose-dependent inhibition of LPA-induced Kenopus oocytes.

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Based on the above results, compound 108 was also examined for its effectiveness as an antagonist of Edg-2, -4, and -7 receptors in RH7777 cells which reterologously express the individual receptors.

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Edg-2, Edg-4, and Edg-7 expressing cells when exposed to a combination of LPA 18:1 Pigure 38 shows the effect of compound 108 on the Ca²⁺ responses in and compound 108. For these experiments, the concentration of LPA was chosen to be near the EC30. Compound 108 significantly inhibited the Ca2+ responses to about 53% and 56% of control, respectively, in Edg-2 and Edg-7 expressing cell lines. In

WO 01/71022 -84 - PCT/U

contrast, compound 108 significantly increased the Ca²⁺ responses to about 148% of control in Edg-4 expressing cell lines.

Therefore, the straight-chain phosphates would be expected to selectively inhibit Edg-2 and Edg-7 activity in vivo and selectively enhance Edg-4 activity in vivo.

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WO 01/71022

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PCT/US01/08729

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Although preferred embodiments have been depicted and described in detail herein, it will be apparent to those skilled in the relevant art that various modifications, additions, substitutions, and the like can be made without departing from the spirit of the invention and these are therefore considered to be within the scope of the invention as defined in the claims which follow.

-92-

What Is Claimed Is:

. A compound according to formula (I)

vherein.

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at least one of X', X', and X' is (HO)₂PO—Z'— or (HO)₂PO—Z'—P(OH)O—Z'—, X' and X' are linked together as —O—PO(OH)—O—, or X' and X' are linked together as —O—PO(OH)—NH—;

at least one of X^1 , X^2 , and X^3 is R^1-Y^1-A- with each being the same or different when two of X^1 , X^2 , and X^3 are R^1-Y^1-A- , or X^2 and X^3 are linked together as $-N(H)-C(O)-N(R^1)-$;

5

optionally, one of X^1, X^2 , and X^3 is H;

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A is either a direct link, (CH₂), with k being an integer from 0

Y' is -(CH₂)- with I being an integer from 1 to 30, -0-,

to 30, or O;

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 Q^1 and Q^2 are independently $H_2 = NR^4$, = 0, a combination of H

 Z^2 is $-(CH_2)_n$ or $-O(CH_2)_n$ with n being an integer from

and -NR'R';

1 to 50 or -0-

23

R¹, for each of X¹, X², or X³, is independently hydrogen, a straight or branched-chain C1 to C30 alkyl, a straight or branched-chain C2 to C30 alkenyl, an aromatic or heteroaromatic ring with or without mono-, di-, or tri-

WO 01/71022

PCT/US01/08729

- 93 -

substitutions of the ring, an acyl including a C1 to C30 alkyl or an aromatic or heteroaromatic ring, an arylalkyl including straight or branched-chain C1 to C30 alkyl, an aryloxyalkyl including straight or branched-chain C1 to C30 alkyl,

$$\begin{array}{c} CH \\ NH \\ NH \\ NH \\ NH \\ R^2 \\ R^8 \\ NR^8 \\$$

R², R³, R⁴, R⁵, R⁶, R⁷, and R⁸ are independently hydrogen, a straight or branched-chain C1 to C30 alkyl, a straight or branched-chain C2 to C30 alkenyl, an aromatic or heteroaromatic ring with or without mono-, di-, or trisubstitutions of the ring, an acyl including a C1 to C30 alkyl or aromatic or heteroaromatic ring, an arylalkyl including straight or branched-chain C1 to C30 alkyl, or an aryloxyalkyl including straight or branched-chain C1 to C30 alkyl;

wherein the compound of formula (f) is not lysophosphatidic acid, phosphatidic acid, cyclic phosphatidic acid, alkenyl glycerolphosphate, dioctyl glycerol pyrophosphate, or N-palmitoyl-L-serine.

The compound according to claim 1, wherein
Q¹ and Q² are both H₂;
one of X¹, X², and X³ is (HO)₂PO—Z²—P(OH)O—Z²—, with

15

 Z^1 and Z^2 being O; and two of X^1 , X^2 , and X^3 are R^1-Y^1-A- , with A being a direct link and Y^1 being O for each.

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4.

PCT/US01/08729

The compound according to claim 1, wherein Q1 is H2;

Q2 is =0;

X2 and X3 are R1-Y1-A-, with A being a direct link and Y1 X1 is (HO)2PO-Z1-, with Z1 being O; and

being -NH- for each.

The compound according to claim 3, wherein X3 is -NH2 and X2 is -NHR1 with R1 being a C14 to C18 alkyl. The compound according to claim 4, wherein R1 is a C14 allcyl.

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The compound according to claim 4, wherein R1 is a C18 alkyl. 6.

X3 is —NHR1 with R1 being an acetyl group and The compound according to claim 3, wherein X2 is -NHR1 with R1 being a C14 alkyl. ζ.

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The compound according to claim 1, wherein Q1 is =NR4; œ

Q2 is H2;

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X3 is R1-Y1-A-, with A being a direct link and Y1 being X1 and X2 are linked together as -0-PO(OH)-0-; and

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The compound according to claim 1, wherein Q1 and Q2 are both H2; ö

two of X1, X2, and X3 are (HO),PO-Z1-, with Z1 being O;

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one of X1, X2, and X3 is R1-Y1-A-, with A being a direct

link and Y1 being --0-

WO 01/71022

PCT/US01/08729

- 95 -

10. The compound according to claim 9, wherein R¹ is an acyl including a C21 alkyl. The compound according to claim 9, wherein R is a C18 alkyl Ξ.

A pharmaceutical composition comprising: a pharmaceutically-acceptable carrier and a compound according to claim 1. 12

A method of inhibiting LPA activity on an LPA receptor 13

comprising

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providing a compound according to claim 1 which has activity as an LPA receptor antagonist and

conditions effective to inhibit LPA-induced activity of the LPA receptor.

2

contacting an LPA receptor with the compound under

 The method according to claim 13, wherein the LPA receptor is present on a cell and said contacting is carried out in vitro. The method according to claim 13, wherein the LPA receptor is present on a cell and said contacting is carried out in vivo. 15.

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The method according to claim 13, wherein the LPA receptor is selected from the group consisting of EDG-2, EDG-4, EDG-7, and PSP-24.

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providing a compound according to claim 1 which has activity A method of modulating LPA receptor activity comprising: contacting an LPA receptor with the compound under as either an LPA receptor agonist or an LPA receptor antagonist and 17.

conditions effective to modulate the activity of the LPA receptor. ဓ

The method according to claim 17, wherein the LPA receptor is present on a cell and said contacting is carried out in vitro.

WO 01/71022 - 96 - PCT/US01/08729

19. The method according to claim 17, wherein the LPA receptor is present on a cell and said contacting is carried out in vivo.

- The method according to claim 17, wherein the LPA receptor is selected from the group consisting of EDG-2, EDG-4, EDG-7, and PSP-24.
- 21. The method according to claim 17, wherein the compound has activity as an LPA receptor agonist and said contacting is carried out under conditions effective to induce LPA receptor activity.

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22. The method according to claim 17, wherein the compound has activity as an LPA receptor antagonist and said contacting is carried out under conditions effective to reduce LPA receptor activity.

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- 23. A method of treating cancer comprising: providing a compound according to claim 1 and administering an effective amount of the compound to a patient in a manner effective to treat cancer.
- 24. The method according to claim 23, wherein the cancer is prostate cancer or ovarian cancer.

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- 25. The method according to claim 23, wherein the compound is an LPA receptor antagonist and said administering comprises:

 delivering the compound to cancer cells, where the compound binds to LPA receptors to inhibit proliferation or metastasis of the cancer cells.
- 26. The method according to claim 23, wherein upon deliverying the compound to cancer cells, the cancer cells are destroyed.

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WO 01/71022 PCT/US01/08729 - 97 -

27. A method of enhancing cell proliferation comprising: providing a compound according to claim I which has activity as an agonist of an LPA receptor and

contacting the LPA receptor on a cell with the compound in a manner effective to enhance LPA receptor-induced proliferation of the cell.

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- 28. The method according to claim 27, wherein the LPA receptor is selected from the group consisting of EDG-2, EDG-4, EDG-7, and PSP-24...
- The method according to claim 27, wherein the cell is in vitro.

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- The method according to claim 27, wherein the cell is in vivo.
- 31. A method of treating a wound comprising:

 providing a compound according to claim I which has activity
 as an agonist of an LPA receptor and

 delivering an effective amount of the compound to a wound site,
 where the compound binds to LPA receptors on cells that promote healing of the
 wound, thereby stimulating LPA receptor agonist-induced cell proliferation to promote
- 32. The method according to claim 31, wherein said delivering comprises:

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wound healing.

introducing to the wound site a composition comprising the compound and a pharmaceutically acceptable carrier.

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33. The method according to claim 32, wherein the wound site is external and said introducing comprises:

topically applying the composition to the wound site.

A method of making a compound according to claim 1

comprising:

reacting (Y²O)₂PO—
$$Z^{11}$$
— Z^{13} or (Y²O)₂PO— Z^{12} —P(OH)O— Z^{11} — Z^{13} ,

$$Z^{11}$$
 is $-(CH_2)_n$ or $-O(CH_2)_n$ with m being an integer

$$Z^{12}$$
 is $-(CH_2)_n$ or $-O(CH_2)_n$ with n being an integer

from 1 to 50 or --0-;

$$Z^{13}$$
 is H or a first leaving group or $-Z^{11}-Z^{13}$ together form the

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$$Z^{13}$$
 is H or a first leaving group or $-Z^{11}-Z^{13}$ together

first leaving group; and

Y2 is H or a protecting group,

with an intermediate compound according to formula (VI)

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at least one of X11, X12, and X13 is R11-Y11-A- with each being the same or different when two of X11, X12, and X13 are R11- Y^{11} —A—, or X^{12} and X^{13} are linked together as —N(H)—C(O)— N(R¹¹)

at least one of X11, X12, and X13 is OH, NH2, SH, or a second leaving group;

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A is either a direct link, (CH₂), with k being an integer from 0

to 30, or O;

23

Q' and Q' are independently H2, =NR13, =O, a combination of

H and -NR 14R15;

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WO 01/71022

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PCT/US01/08729

straight or branched-chain C1 to C30 alkyl, a straight or branched-chain C30 alkyl or an aromatic or heteroaromatic ring, an arylalkyl including C2 to C30 alkenyl, an aromatic or heteroaromatic ring with or without R11, for each of X11, X12, or X13, is independently hydrogen, a straight or branched-chain C1 to C30 alkyl, an aryloxyalkyl including mono-, di-, or tri-substitutions of the ring, an acyl including a Cl to straight or branched-chain C1 to C30 alkyl,

straight or branched-chain C1 to C30 alkyl, a straight or branched-chain C2 to C30 alkenyl, an aromatic or heteroaromatic ring with or without R12, R13, R14, R15, R16, and R17 are independently hydrogen, a C30 alkyl or aromatic or heteroaromatic ring, an arylalkyl including mono-, di-, or tri-substitutions of the ring, an acyl including a C1 to straight or branched-chain C1 to C30 alkyl, or an aryloxyalkyl including straight or branched-chain C1 to C30 alkyl;

according to formula (f) where one or two of X^1, X^2 , and X^3 is (HO)₂PO— Z^1 — or deprotection step being performed under conditions effective to afford a compound followed by a de-protection step, if necessary, with both said reacting and the (HO),PO-Z2-P(OH)O-Z1-.



PCT/US01/08729

PCT/US01/08729

1/26

PSP 24 Family

6

GPM

GPM

EDG Family -- Edg 1 SPM Edg 2 GPM Edg 7 GPM Edg 3 SPM Edg 4 GPM Edg 5 SPM Edg 6 SPM Edg 8 SPM

Figure 1

∱Boo-L-Serine 24 TFA 2/26 · *** 26-34

Figure 2

26 R = (CH₂)₃CH₃
27 R = (CH₂)₁₃CH₃
28 R = (CH₂)₁₇CH₃
29 R = PC₆H₂O(CH₂)₁₃CH₃
30 R = PC₆H₂O(CH₂)₁₃CH₃
31 R = m-C₆H₂O(CH₂)₁₃CH₃
32 R = m-C₆H₄O(CH₂)₁₃CH₃
33 R = c-C₆H₄O(CH₂)₁₃CH₃
34 R = c-C₆H₄OCH₃

36 R = (CH₂)₆CH₃
36 R = (CH₂)₁₃CH₃
37 R = (CH₂)₁₇CH₃
38 R = p-C₆H₄O(CH₂)₁₃CH₃
39 R = p-C₆H₄O(CH₂)₁₃CH₃
40 R = m-C₆H₄O(CH₂)₁₃CH₃
41 R = m-C₆H₄O(CH₂)₁₃CH₃
42 R = o-C₆H₄O(CH₂)₁₃CH₃
43 R = o-C₆H₄O(CH₂)₁₃CH₃

PCT/US01/08729

PCT/US01/08729

WO 01/71022

Figure 3

63 R = (CH₂)₁₇CH₃ 64 R = (CH₂)₁₁CH₃ 65 R = (CH₂)₁₅CH₃ Tetrazole
 Dibenzyldlisopropyl
 phosphoramidate 4/26 3) Peracetic acid

60 R = (CH₂)₁₇CH₃ 61 R = (CH₂)₁₁CH₃ 62 R = (CH₂)₁₆CH₃

Pd-C/H₂

66 R = (CH₂)₁₇CH₃ 67 R = (CH₂)₁₁CH₃ 68 R = (CH₂)₁₆CH₃

Figure 4

PCT/US01/18729

6/26

5/26

Tebazole
 Dibenzyldilsopropyl
 phosphoramidate

Peracetic acid

69-76

77-84

77 R = (CH₂)₁₂CH₃
78 R = (CH₂)₁₃CH₃
79 R = (CH₂)₁₄CH₃
80 R = (CH₂)₁₆CH₃
81 R = (CH₂)₁₆CH₃
82 R = (CH₂)₁₇CH₃
83 R = (CH₂)₁₈CH₃
84 R = (CH₂)₂₀CH₃

69 R = (CH₂)₁₂CH₃
70 R = (CH₂)₁₃CH₃
71 R = (CH₂)₁₄CH₃
72 R = (CH₂)₁₆CH₃
73 R = (CH₂)₁₆CH₃
74 R = (CH₂)₁₆CH₃
76 R = (CH₂)₁₆CH₃
76 R = (CH₂)₁₆CH₃

71.84

86 R = (CH₂)₁₂CH₃
86 R = (CH₂)₁₃CH₃
87 R = (CH₂)₁₄CH₃
88 R = (CH₂)₁₅CH₃
89 R = (CH₂)₁₆CH₃
90 R = (CH₂)₁₆CH₃
91 R = (CH₂)₁₆CH₃
92 R = (CH₂)₁₆CH₃

Figure 5A

Figure 5B

III) Peracetic acid N) 10% Pd/C, H₂

Figure 6A

7/26

C₆H₆CH₂OH/(CF₈SO₂)₂O,

ROH BF, CHC,

KO₂, 18-crown-8 DMF, DME, DMSO

Pd(OH)₂/C, H₂ CH₃OH, H₂O

Pyridine, p-TsCl

Figure 7C

Figure 6B

Figure 11

III) Peracetto acid

Figure 12

WO 01/71022

PCT/US01/08729

10/26

Figure 10

11/26

WO 01/71022

PCT/US01/08729

Figure 13

1. Tris(1,2,4-triazole)phosphate 2. 2% HCi

HO NH, T-R NH, T-COO-

36-43

Figure 14

50-64

10% Pd/C, H₂

2) Peracetic Acid

PCT/US01/08729

2. H₂/Pd/C

Figure 19

WO 01/71022

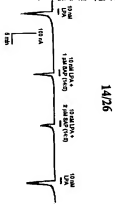


Figure 21

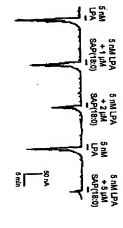


Figure 22

Figures 23A-B

Figure 20

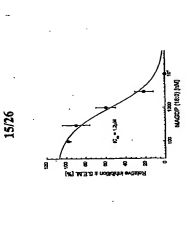
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10 mM.PA + 2 µM.6DAP (14:0/2:0)

اِلْمَارِ الْمَارِ Figure 26





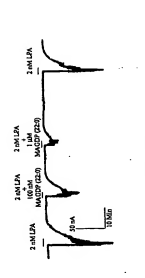


Figure 25

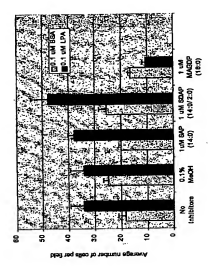
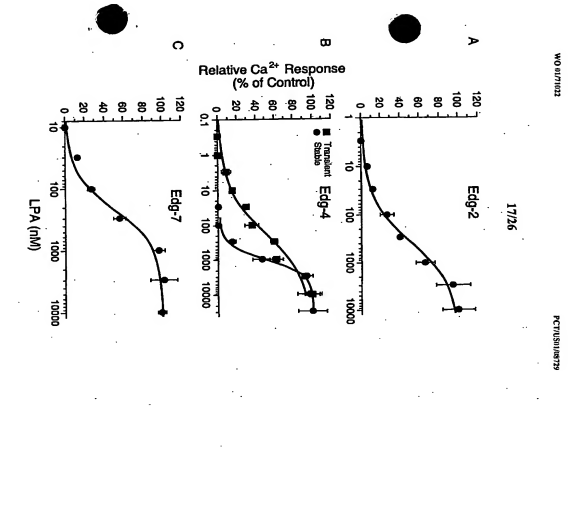


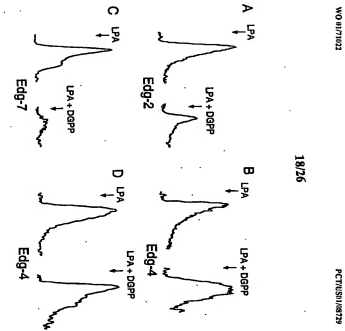
Figure 27

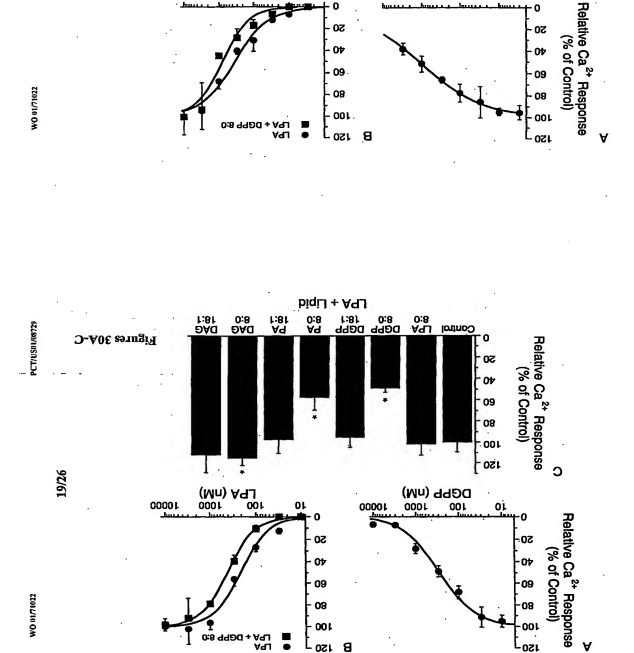
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Figures 28A-C

Figures 29A-D





B 150

biqiJ + AqJ

ÐA₫ 0:8

DAG 1:81

(Mn) Aqj

10

100 1000 10000

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D-Alf esrugiA

A9 1:81

Control LPA DGPP DGPP 1:81 0:8 0:8

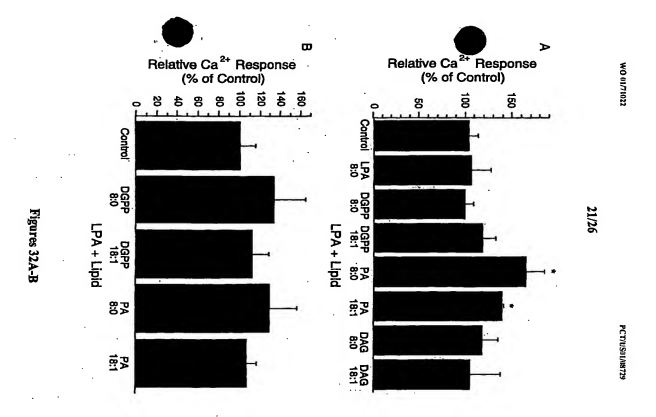
(Mn) 9920

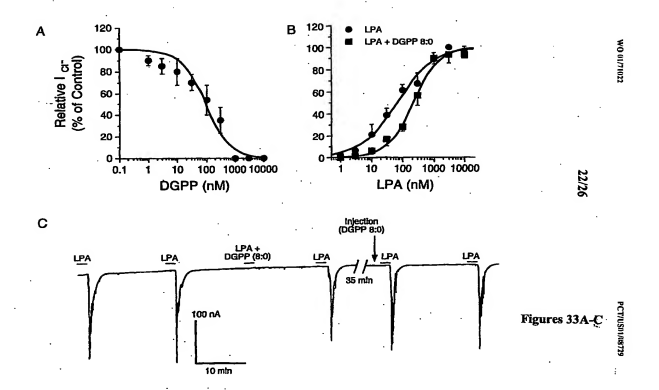
1000 10000 100000

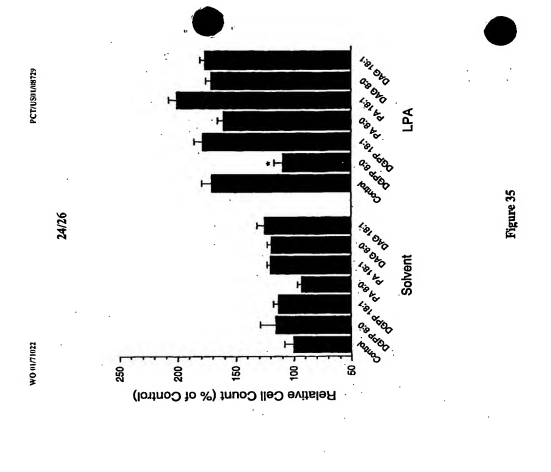
Relative Ca²⁺ Response (% of Control)

၁

-09 08 100







Control DGPP DGPP PA 8:0 8:181 8:0 Diqil + Aql

LPA+ LPA+ S1P + S1P + Solvent DGPP

A9 1:81

PCT/US01/IB729

WO 01/71022

23/26

Figures 34A-D

Relative Ca²⁺ Response (% of Control)

Relative Ca²⁺ Response (% of Control)

·8

a

- oż

-07

-001 -08 -09

Lost

20

01

150

PSP24

PSP24

12345678

15348678

:6p∃

:6p∃

— 25.0 (dd)

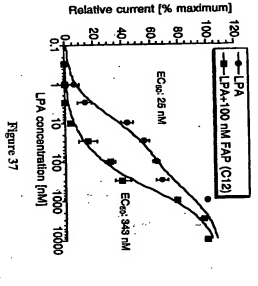
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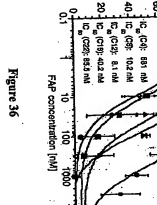
(pq)

O.25 —

PCT/US01/08729

-FAP (C4) -FAP (C8) -FAP (C12)





Relative inhibition [% control]

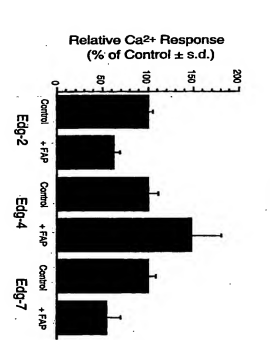


Figure 38

PCT/US01/IB729 WO 01/71022

SEQUENCE LISTING

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<210> 1 <211> 1095

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<210> 2 <211> 364

WO 01/71022

PCT/US01/08729

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Leu Ala Asn Lou Lou Val Met Val Ala Ile Tyr Val Asn Arg Arg Pho 65

His Phe Pro Ile Tyr Tyr Leu Met Ale Ann Leu Ala Ala Ala Asp Phe 95

Phe Ala Gly Leu Ala Tyr Phe Tyr Leu Mot Phe Asn Thr Gly Pro Asn 100

The Arg Arg Leu The Val Ser The Trp Lou Lou Arg Gln Gly Leu Ile 115

Asp Thr Ser Leu Thr Ala Ser Val Ala Asn Leu Leu Ala Ile Ala Ile

Glu Arg His Ile Thr Val Phe Arg Met Gln Leu His Thr Arg Met Ser 150 145 Asn Arg Arg Val Val Val Ile Val Val Ile Trp Thr Met Ala Ile 175 val Met Gly Ala Ile Pro Ser Val Gly Trp Asn Cys Ile Cys Asp Ile

Glu Asn Cys Sar Asn Met Ala Pro Leu Tyr Ser Asp Ser Tyr Leu Val

Phe Trp Ala Ile Phe Asn Leu Val Thr Phe Val Val Met Val Val Leu

Tyr Ala His Ile Phe Gly Tyr Val Arg Gln Arg Thr Met Arg Met Ser 225

WO 01/71022 PCT/US01/08729

Į eu Arg His Sor Ser Gly Pro Arg Arg Asn Arg Asp Thr Met Met Ser Leu 245 250 255 Lys Thr Val Val Ile Val Lou Gly Ala Phe Ile Ile Cys Trp Thr 260 270

Pro Gly Lou Val Leu Leu Leu Asp Val Cys Cys Pro Gln Cys Asp 275 280 285

Val Leu Ala Tyr Glu Lys Phe Phe Leu Leu Leu Ala Glu Phe Asn Ser 290 295

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325 330 Ala Met Asn Pro Ile Ile Tyr Ser Tyr Arg Asp Lys Glu Met Ser Ala 305 310 320

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> WO 01/71022 PCT/US01/08729

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Thr Asn Leu Leu Val Ile Ala Ala Ile Ala Ser Asn Arg Arg Pho His 50 55

Gln Pro Ile Tyr Tyr Leu Leu Gly Asn Leu Ala Ala Ala Asp Leu Phe 65 70 75

Ala Gly Val Ala Tyr Leu Phe Leu Met Phe His Thr Gly Pro Arg Thr 90 95 95

Ala Arg Leu Ser Leu Glu Gly Trp Phe Leu Arg Gln Gly Leu Leu Asp 100 105 The Ser Leu The Ale Ser Vel Ale The Leu Leu Ale Ile Ale Vel Glu
115 120 125

Αřg His Arg Ser Val Met Ale Vel Gln Leu Hie Ser Arg Leu Pro Arg 130

Gly Arg Val Val Mot Lou Ile Val Gly Val Trp Val Ala Ala Lou 145 150 G1y 160

Leu Gly Leu Leu Pro Ala His Ser Trp 165 His Cys Leu Cys Ala Leu 170 175 Asp

Arg Cys Ser Arg Met Ala Pro Leu Leu Ser Arg Ser Tyr Leu Ala Val 180

Trp Ala Leu Ser Ser Leu Leu Val Phe Leu Leu Met Val Ale Val Tyr 195 200 205

PCT/US01/II8729 WO 01/71022

Thr Arg 11e Phe Phe Tyr Val Arg Arg Arg Val Gln Arg Met Ala Glu 210 His Val Ser Cys His Pro Arg Tyr Arg Glu Thr Thr Leu Ser Leu Val 225 Lys Thr Val Val Ile Ile Leu Gly Ala Phe Val Val Cys Trp Thr Pro 250 250 Val Leu Leu Leu Asp Gly Leu Gly Cys Glu Ser Cys Asn 260 Leu Ala Val Glu Lys Tyr Phe Leu Leu Leu Ala Glu Ala Ash Ser Leu Val Asn Ala Ala Val Tyr Ser Cys Arg Asp Ala Glu Met Arg Arg Thr Phe Arg Arg Leu Leu Cys Cys Ala Cys Leu Arg Gln Ser Thr Arg 305 Glu Ser Val His Tyr Thr Ser Ser Ala Gln Gly Gly Ala Ser Thr Arg Pro Glu Asn Gly His Pro Leu Met Asp Ser Thr Leu 340 350 295 Gln Val 275 Ile Met Leu G1, Val

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WO 01/71022

PCT/US01/08729

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Val Leu Cys Val Gly Thr Phe Phe Cys Leu Phe Ile Phe Phe Ser Asn

Phe Tyr Tyr Leu Leu Ala Asn Leu Ala Ala Ala Asp Phe Phe Ala Gly 5

Ile Ala Tyr Val Phe Leu Met Phe Asn Thr Gly Pro Val Ser Lys Thr 95

Leu Thr Val Asn Arg Trp Phe Leu Arg Gln Gly Leu Leu Asp Ser Sor 105 Leu Thr Ala Ser Leu Thr Asn Leu Leu Val Ile Ala Val Glu Arg His 126

Val Thr Leu Leu Ile Leu Leu Val Trp Ala Ile Ala Ile Phe Met Gly 145 Met Arg Val His Ser Asn Leu 135 130

Lys Lys Arg

뵱

Ser Ile Met Arg

Ala Val Pro Thr Leu Gly Trp Asn Cys Leu Cys Asn Ile Ser Ala Cys
175

Ser Ser Leu Ala Pro Ile Tyr Ser Arg Ser Tyr Leu Val Phe Trp Thr 190 185 180

WO 01/71022 PCT/US01/08729

Val Mot Thr Val Lou Gly Ala Pho Val Val Cya Trp Thr Pro Gly Lou 250 255 Ile Val Ser Asn Leu Met Ala Phe Leu Ile Met Val Val Val Tyr Leu Arg Gly Ser Ile Ser Arg Arg Arg Thr Pro Met Lys Leu Met Lys Thr 230 236 TY. ΔĐ Tyr Val Lys Arg Lys Thr Asn Val Leu Ser Pro 215 220 200 205 His Thi

Val Val Lou Lou Asp Gly Lou Asn Cys Arg Gln Cys Gly Val Gln 265 270

His Val Lys Arg Trp Pho Leu Leu Leu Ala Lou Leu Asn Ser Val Val 275 280 285

Lys Lys Mot Ile Cys Cys Pho Ser Gln Glu Asn Pro Glu Arg 305 310 315 Pro Ile Ile Tyr Ser Tyr Lys Asp Glu Asp Met Tyr Gly Thr Met 290 295 300 βīγ 920

Agn

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Thr Thr Phe Val Val Tyr Glu Aen Thr Tyr Met Aen Ile Thr Leu 20 25 30

Ala Val Pro 65 Thr Thr Pro Ala Ala Phe Lys Ser Leu Asn Leu Pro 70 75 90 Ten

Gln Ile Thr Leu Ser Ala Ile Met Ile 85 Phe Ile Leu Phe Val Ser 90 95 양성

Arg neg Ser Ala Ile Asn Ile Leu Leu Ala Ser Leu Ala Phe Ala Asp Met 115 120 125 . Gly Asn Leu Val Val Cys Leu Met Val Tyr Gln Lys Ala Ala Met 100 105

Leu Leu Ala Val Leu Asn Met Pro Phe Ala Leu Val Thr Ile Leu 135 140

PCT/US01/08729

WO 01/71022

WO 01/71022	
PCT/US01/08729	
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PCT/US01/08729

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Met	175	Asn	Phe		Ţ	Phe 255	H1.8	Gln	391	Let		Glu Ile	ng.	203	11.0
Ala	110	Leu 190	Ser Phe	118	G1.y	Pro	Arg 270	Ser	ig t	119	Ser Leu 335	Phe (ne.	873	ily l
Ser	Leu	Lys	7 205	Gln Ile Pro	Pro	Phe Phe Ile Pro	reg T	Leu 285	31n l	Thr Ile Leu Ile Leu Phe 315	ľyr :	Tyr Gln His Asn Phe Phe 345	Ala 1 365	Ala (Gln Leu Pro Gly His Thr 395 '400
Val	Leu	Asp		Leu 220	Agn	Phe	Ę	Cys	Phe 300	[]e	Ę	· §	Ser /	Asp 1	ne n
Arg. 155	118	gln	Ę.	Zg Zg	Thr. 235	Phe	- Ren	119	or c	Thr 315	Ę	118	ys .	11.8	Gln 1 395
S,	A18 170	Arg	Ser	Pro	Į	Ser 250	neg 7	ily .	LE 1		Phe 1	ı.	1 ne	e d	2
Phe	Za Z	Gln 185	Za1	- Ren	ž	3	11e) 265	37n (ll 1	be 1	2	Tyr (ž.	ξ.	n n
å	31.y	/a1	Ala 200	17 7	, t	, e	71	Pro (280	S	L'a	1 81	¥	Cys 1 360	ya I	he I
. eya	ll (116		Val (215	e d	er 1	et O	74	Ser Leu Gln Arg Pro Phe Gln Met 295	6	ĝ.	r er	9 8	Ile 1 375	8
Gly 1 150	9	[B	. ne.	La 1	Val 1]e	e c	Ä		Thr A	ž.	8 1	E	ב ה ה	Phe L 390
9,	val 1	19 ₀	'a1 1	20 7	7 8 %	Leu 1 245	E I	8 8	a a	78 1	Val C 325	E SA		e e	A E
The Arg Trp Ile Phe Gly Lys Phe Phe Cys Arg Val Ser Ala Met 145	Phe Trp Leu Phe Val Ile Glu Gly Val Ala Ile Leu Leu Ile Ile 165	ile Asp Arg Phe Leu ile Ile Val Gin Arg Gin Asp Lys Leu Asn Pro 185	Ale Lys Vel Leu Ile Ala Vel Ser Trp Ale 195	Ala Phe Pro Leu Ala Val Gly Asn Pro Asp Leu 210	Arg Ala Fro Gln Cys Val Fhe Gly Tyr Thr Thr Asn Pro Gly Tyr 225	ile Leu ile Ser Leu ile Ser 245 250	Val ile Leu fyr Ser Phe Met Gly Ile Leu Asn Thr Leu Arg His Asn 265 270	Ala Leu Arg Ile His Ser Tyr Pro Glu Gly Ile Cys Leu Ser Gin Ala 275	Ser Lys Leu Gly Leu Met 290	Asp Met Gly Phe Lys Thr Arg Ala Phe Thr 305	Ala Val Pho Ile Val Cys Trp Ala Pro Phe Thr Thr Tyr 325	Ala Thr Phe Ser Lys His Phe Tyr 340	Sor Thr Trp Leu Leu Trp Leu Cys Tyr Leu Lys Sor Ala Leu Asn Pro 355	Leu Ile Tyr Tyr Trp Arg Ile Lys Lys Phe His Asp Ala Cys Leu 370	Met Met Pro Lys Ser Phe Lys Phe Leu Pro 395
E E	و . م	F T	Ala L 195	8 6	9 2		2 2	Arg I. 275	9 0	14 P.	e E	ش ش ش	15p Lo 355	7	5 5
T B	17 (2	Q.	4 67 1	ALA P	Ja P	Ala Tyr val	1 e	8u A 2	Lys L 290	ال 9	i i	H H	a a a	11e T3 370	r E
Thr A.	₽ 1	e A	Tyr Arg	Val AU	Arg AJ 225	La T.	H	ž Ž	7. 12.52	υς Σ	ei.	E .	설 성	5 11 E	ή. Σ
F A	Ď.	H	Ĥ.	s'	2 2	4	5	~	ത്	Asp 305	2	2	ល័	, 3	Met 385

ctgctcagcc getcctattt g 4210 16 4211 22 4212 DNA 4212 DNA 4212 DNA 4213 Artificial Sequence: primer, forward EDG-5 4400 16 4213 Dascription of Artificial Sequence: primer, 4212 DNA 4213 Artificial Sequence 4220 42210 DNA 4213 DNA 421	<400> 15	<220> <220> <223> Description of Artificial Sequence: primer, forward EDG-4	<210> 15 <211> 21 <212> DNA	<pre><213 Artificial Sequence <213 Artificial Sequence <220> <222> Description of Artificial Sequence: primer, reverse EDG-3 <400> 14 tgctgstgca gaaggcaatg ta</pre>	<400> 13 cttggtcatc tgcagcttca tc <210> 14 <211> 22	<210> 13 <211> 22 <212> DNA <213> Artificial Sequence <220> <220> <223> Description of Artificial Sequence: primer, forward EDG-3	<220> <223> Description of Artificial Sequence: primar, reverse EDG-2 <400> 12 gttggccatc asgtastass ta	<211> 22 <212> DNA <213> Artificial Sequence
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12

WO 01/71022

PCT/US01/08729

WO 01/71022

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WO 01/71022	PCT/US01/IN729	WO 01/71022	PCT/US01/08729
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Tamas: 57 N. Somerville #401, Memphis, TN 38104 (US). NUSSER, Nora: 57 N. Somerville #213, Memphis, TN 38104 (US).

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(54) Title: LPA RECEPTOR AGONISTS AND ANTAGONISTS AND METHODS OF USE

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(57) Abstract: The present invention relates to compounds exceeding to formula () as disclosed herein as well as pharmaceutical compounds which include those compounds. Also disclosed are methods of using such compounds, which have earlyly as agonists of LPA receptors; such methods including inhibiting LPA activity on an LPA receptor, modulating LPA receptor, modulating LPA receptor.

INTERNATIONAL SEARCH REPORT

International application No.

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Form PCT/ISA/210 (second sheet) (July 1998) •

INTERNATIONAL SEARCH REPORT

International application No.	PCT/US01/(1K729	
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×i<	Database Caplus on STN, Chemical Abstracts (Columbus Ohio, USA), CA:87:33513, GIBBS, D. 'The synthesis of phosphoramidates from silylphosphites and azides', abst Tetrahedron Lett. (8), pages 679-82, 1977.	1.5.7.
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A,X	Database Caplus on STN, Chemical Abstracts (Columbus Ohio, USA), CA:134:174618, BADALASSI, F. 'A versatile periodate-toupled fluorogenic assay for hydrolytic enzymes' abst Angew. Chem. Int Ed. 39(22) pages 4067-4070, 2000.	1, 9
×	Database Caplus on STN, Chemical Abstracts (Columbus Ohio, USA), CA:99:3425, BUSHNEV. A.S. et al. 'Synthesis of rac-3-benzoyl-1-deoxyceramide-1-phosphonic acid' abst Bioorg. Khim 9 (4) pages 533-5, 1983.	-
×	Database Caplus on STN Chemical Abstracts (Columbus Ohio, USA), CA:75:71703, CATES, L. 'Phosphorus-nitrogen compounds . 12. Phosphamidase studies. 2. N-alkylphosphoramidic acids' abs J. Med. Chem. 14(7) pages 647-9, 1971.	1, 12
×	Database Caplus on STN, Chemical Abstracts (Columbus Ohio, USA), CA:116:221452, GAS CO, M.R. et al 'Timolol in lipospheres' abst Pharmazie 47 (2) pages 119-21, 1992.	1, 12
×	Database Caplus on STN, Chemical Abstracts (Columbus Ohio, USA), CA:76:34548, AVAEVA, S. M. et al 'Hydroylsis of phosphoric ester serine derivatives containing free anino or carboxy groups' abst Vestn. Mosk. Univ. Khim. 12(5) pages 627-8, 1971.	1, 3
×i«	Database Caplus on STN, Chemical Abstracts (Columbus Ohio, USA), CA:116:236128, TONG G. et al 'Synthesis of the simple peptide model Ac-Abu(PO3H2)-NHMe' abstr Aust. J. Chem. 45(4) pages 777-84, 1992.	1, 3

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INTERNATIONAL SEARCH REPORT

	International application No. PCT/US01/08729
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J. DOCUMENTS CONSIDERED TO BE RELEVANT Challon of document, with indication, where appropriate, of the relevant passages	Database Caplus on STN, Chemical Abstracts (Columbus Ohio, USA), CA:126:16188, RYAN, M. et al 'Synthesis, structure-activity relationships, and the effect of polyethylene gycol on inhibitors of phosphatidylinositol specific phospholipase C from Bacillus cereus, abst J. Med. Chem. 39(22) pages 4366-4376, 1996.	
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International application No. PCT/US01/08729

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Remark on Protest			_	This International Searching Authority found multiple inventions in this international application, as follows: Please See Estan Sheet.	Box II	ľ		_	This insernational report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:	Be .
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7	urici e	and J-7	ching.	Sc	13			i i	E	3
1	No required additional search fees were timely paid by the applicant. Consequently, restricted to the invention first mornimed in the claims; it is covered by claims Nos.:	As all rearchable claims could be rearched without effort justifying an additional fee, this Authority did not invite payment of any additional fee. As only arome of the required additional search fees were timely paid by the applicant, this international search report cover only those claims for which fees were paid, specifically claims Nos.: 1.3.7-and 9-12 (in pan)	As all required additional scarch feer were timely paid by the applicans, this international search report covers all scarchable chims.	xemational Searching / Please See Extra Sheet	Observations where unity of invention is lucking (Continuation of item 2 of first abost	Claims Nos.: beause they are dependent chims and are not draited in accordance with the second and third septences of Rule 6.4(s)	Claims Nov.: because they relate to parts of the international application that do not comply with the prescribed requirements to nuch an extent that no meaningful international search cun be carried out, specifically:	Chilin Non: because they relate to subject matter not required to be searched by this Authority, namely:	g g	Observations where certain claims were found unsurchable (Continuation of item 1 of first sheet)
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INTERNATIONAL SEARCH REPORT

International application No. PCT/US01/08729

A. CLASSIFICATION OF SUBJECT MATTER: US CL :

562/ 8. 11. 12, 15, 20, 23, 24; 546/22; 548/112; 549/5, 6, 7, 8, 9, 10, 11, 12, 13, 218

B. FIELDS SEARCHED

Electronic data bases consulted (Name of data base and where practicable terms used):

EAST, CAS ON LINE, BEILSTEIN

search terus: structure drawing, monophosphate ester, phusphate amide ester

This ISA found multiple inventions as follows: BOX II. OBSERVATIONS WHERE UNITY OF INVENTION WAS LACKING

This application consists chims directed to more than one invention. These are deemed to back Unity of Invention because they are not so inteed as to form a single inventive concept under PCT Rule 13.1 sizes they have no special technical feature which unless them.

Group I, claims 1-2(in part), 9-12(in part), drawn to compounds of formula (f) wherein there is no cyclic ring structure which contains a linking of X1 and X2 or of X2 and X3 and at least one of X1 or X2 or X3 is R1-Y1-A. wherein R1 is not a heterocyclic ring or or a disregar containing group and Y1 is (CH2), or O or C(O), A is a link or (CH2), or O and Z1 is O (CH2), or O . 22 is O or O(CH2)m. Q1 and Q2 are H2 or O. (i.e. monophosphase enter-(HO),P(O)-O-CQ1— or linked enter-PO-Z2-P)

Q1.--). Group II. (skinds) (lin part), 12(in part), driven to compounds of formula (f) wherein there is no cyclic ring structure which contains a linking of X1 and X2 or X2 and X3 and x1 least one of X1 or X2 or X3 is R1-V1-A wherein R1 is not a heterosylic finite or or a stringen containing group and Y1 is (CH2), or O. or carbonyl. A is a link or (CH3), or O. or carbonyl. A is a link or (CH3), or O. or CH2 is O. or O(CH2)m. OI and Q2 are H2 or O. (i.e. monophosphorus amide enter -(HO), P(O)NH-

Group III. châmely I (in parth, 12 (in parth, drawn to compounds of the formula (i) wherein there is no cyclic structure which counting a linking of X1 and X2 or of X2 and X3 and at least one of X1 or X2 or X3 is R1.71-A. wherein R1 or not a heterocyclic map or or a random commissing group and Y1 in (CH2), or O. or carbonyl. A is a list or (CH2) or O and Z1 is S, Z2 is O or O(CH2)m. Q1 and Q2 are H2 or O. (i.e. momentisphosphase exter-(HO), p(O)-S-CQ1--).

Group IV, claim(s) 1(in part), 12(in part), drawn to compounds of the control of

Group V. chims 1,37 (in part), 12(in part), drawn to compounds of formula (!) wherein there is no cyclic day arrange with contains a linking of X1 and X2 or of X2 and X2 and at least one of X1 or X2 or X2 is R1.Y1.A wherein R1 is not a latestropylet finight or or a integran containing groups and Y1 is NR2, A is a link or (CH2), or O and Z1 is O (CH2), or O . Z2 is O or O(CH2)m, O1 and Q2 are H2 or O.

Group VI. chint(s): 1.3-7(in pars). 12(in pars), drawn to compounds of formula (1) wherein there is no cyclic ring structure which contains inhibits of XI and X2 or of X2 and X3 and at least one of XI or X2 or X3 is R1-Y1-A wherein R1 is not a heterocyclic rings or or a singular normaling group and Y1 is NR2. A is a link or (CH2), or O and Z1 is NR1. Z2 is 0 or O(CH2)m, Q1 and Q2 are H2 or O. (i.e. monophesphorus smide exter

Group VII, claim(s) I(in part), I2 (in part), dawn to compounds of the formula (I) wherein there is no cyclic ring structure which centains a linking of XI and X2 or of X2 and X3 and at teast one of XI or X2 or X3 is RI-YI-A wherein RI is not a benetocyclic ring or or a rincygen containing wherein RI is not a benetocyclic ring or or a rincygen containing group and YI is NR2. A is a link or (CH2), or O and ZI is S, Z2

Form PCT/ISA/210 (exam sheet) (July 1998) *

REPORT INTERNATIONAL SEARCH

Inemational application No.

PCT/US01/08729

is O or O(CH2)m, Q1 and Q2 are H2 or O. (i.e. monothiophosphasis

-(HO),P(O)-S-CQ1--).

Group VIII, cisten(s) 1(in part), 12(in part), drawn to compounds of the formula (f) wherein there is no cyclic ring entreature which contains a linking of X1 and X2 or of X2 and X3 and at least one of X1 or X2 or X3 in R1-Y1-A wherein R1 is no as the theorycelle fing on or a nitrogen—constituing group and Y1 in KR2, or exhony1, A is a link or (CH3), or O and Z1 is (CH3), or GA3H, Z2 is O or O(CH3)m. Q1 and Q2 arth2 or O. (i.e. phoephorus acid not an enter-(HO),P(O)-CH2-CQ1---).

Group IX, claims 1 (in part), 12(in part), drawn to compounds of formulb (i) wherein there is no cyclis ring structure which comains a linking of X1 and X2 or of X2 and X3 and at least one of X1 or X2 or X3 is R1-Y1-A wherein R1 is not a boterocyclic

ning or or a mitrogen conduining group and Y1 is 10R2, A is a link or (CHI), or O and Z1 is O (CHI), or O . Z2 is O or O(CHI),m,Q1 and Q2 are =NR*-NR?R* or at least one Q1 or Q2 is =NR*-NR?R*.

Group X. claim(s) ((in part). 12(in part). drawn to compounds of formula (t) wherein there is no cyclic ning structure which central and labeling of X1 and X2 or of X2 and X2 and a Lisax or of X1 or X2 or X2 is B1-V4.4 stretch R1 is not a haterogetic ring or or at introgen counting group and Y1 is NR2. A is a lint or (CHZ), or O and Z1 is NH, Z2 is 0 or O(CHZ)m,Q1 and Q2 are = NR* -NR*R* or at least one Q1 or Q2 is = NR* -NR*R* (i.e. monophanphorus amide exer -(HO),P(O)NH-CQ1---).

Group XI. chim(s) I (in part), 12 (in part), 4nwn tu compounds of the formula (i) wharete there is no cyclic ring ancurant which constitus a limiting of XI and X2 or of X2 and X3 and at least one of XI or X2 or X3 is R1-Y1-A wherete R1 is not a text-recycle into or a sincepart consulting.

group and Y1 is NR2. A is a lab or (CH3), or 0 and Z1 is 5, Z
is 0 or Q(CH2) in Q1 and Q2 are =NR* -NR* or at least one Q1 or Q2 is =NR* -NR*R* (i.e. monothlophosphal

ener -(HO),P(O)-S-CQ1---).

Group XII, chim(s) I(in part), 12(in part), drawn to compounds of the formula (I) wherein there is no cyclic ring intrustrue which constants a linking of XI and X2 or of X2 and X3 and at least one of XI or X2 or X2 or X2 is R1-Y1-A wherein R1 is not a heterocycle find or or a subcoping couplaining group and Y1 is NR2, or exabory! A is a link or (CHX), or 0 and Z1 is (CHX), or 0 cRXX+1X is 0 or 0(CHX), m. 0 it and 0.2 old and 0.2 are = NR3-NR3R* (i.e. phosphorus scial not an exter -(HO),P(O.CH2-CQ1---).

Group XIII. chims I (fin part). 12(in part), drawn to compounds of formula (I) wherein there is a cyclic ring struature which contains a linking of XI and X2 into O-PO(OH)-O or X2 and X3 into OPO(OH)-Wit- and at least one of XI or X2 or X1 of XI-X4 wherein R1 is not a heterospile ring or as a introgen containing group and YI is (CH2), or O or C(O). A is a lake or (CH2), or O and Z1 is O (CH3), or O. . Z2 is O or O(CH3). Or or an opphagphase sets-(HO), P(O)-O-CQ1-- or linked exter PO-Z2-P)

Group XIV , claim(s) (its part), 13(in part) , drawn to compounds of formula (i) wherein there is a cyclic ring structure which ordatin's plicibing of XI and XI into CPO(CH)-10 or XI and XI into CPO(CH)-10. Fart at it learn one of XI or XI or XI at its and XI into a sherroxypile ring or or a ninegan containing group and YI is (CH2), or O, or carboxyl , A is a link or (CH2), or O and ZI is NH. ZI is O or O(CH2)m. QI and QI are H3 or O. (i.e. monophosphorus amido exter -(HO),P(O)NH-CQ1---).

Group XV, claim(s) 1(to part). 12 (in part). drawn to compounds of the formula (t) which contains a linking of X1 and X2 are O-PO(O)-PO or X2 and X1 and X2 or X3 is R1-Y1-A wherein R1 x2 are O-PO(O)-PO or X2 and X3 is R1-Y1-A wherein R1 is one a heteropelic fing or a nitrogen containing group and Y1 is (CH2), are O. or carbony).

A is a link or (CH2), or O and Z1 is S. Z2 is O or O(CH2)m. Q1 and Q2 are H2 or O (i.e. monothisphosphate caer •(HO),P(O)-S-CQ1---).

Group XVI. claim(s) 1(in part), 12(in part), dawn to compounds of the formula (1) which constitute (1) which constitute a linking of X1 and X2 into O-PO(OH)-O or X2 and X3 into OPO(OH)-NH- and at least

Form PCT/ISA/210 (extra sheet) (July 1998)

INTERNATIONAL SEARCH REPORT

International application No. PCT/US01/08729 one of XI or X2 or X3 is RI-YI-A wherein RI is nat a heterocyclic ring or or a nitrogen containing group and YI is (CH2), or O or or carboay! A is a link or (CH2), or O and ZI is (CH3), or CR3H, 22 is O or O(CH2)m, QI and Q2 artH2 or O. (i.e. phosphorus acid nas as exter -(HO),P(O-CH2-CQ1--).

Group XVII, chims I (lin purt), 12(in purt), driwn to compounds of formula (f) which corrains a linking of XI and XZ isto OPO(OH)-O or XZ and XZ into OPO(OH)-NH- and at least one of XI or XZ or XZ is is RI-YI-A wherein RI is not a interspecie fing or or a nitrogen containing group and YI is NRZ. A is a link or (CHZ), or O and ZI is O (CHZ)_{or} or O . ZZ is O or O(CHZ)m. Q1 and Q2 are HZ or O.

Group XVIII; claim(s) 1(in part). I3(in part), drawn to compounds of formule (1)which contains a linking of XI and XZ into O-PO(OR)-O or XZ and XZ into OPO(OR)-NH. and at least one of XI or XZ or XZ in IR-YI-A wherein RI is not a heteropytic fing or or a mirogen commulating group and YI is NR2, A is a link or (CH2), or 0 and ZI is NR, ZZ is O or O(CH2)m, QI and QZ are HZ or O. (i.e. monophaphorus amide exer-(HO), P(O)NH-CQ1-...).

Group XIX, chain(s) 1(in part), 12 (in part), drawn to compounds of the formula (i) which contains a linking of X1 and X2 into O-PO(OH)-O or X2 and X3 into O-PO(OH)-WI- structure and at least one of X1 or X2 or X3 is R1-Y1-A wherein R1 is not a heterospaic ring or or a nitrogen constituing group and Y1 is NR2 . A is a link or (CH2) , or O and 21 is S. Z2 is 0 or O(CH2)m, Q1 and Q2 are H2 or O. (i.e. monothirp/hosphase cner -(HO),P(O)-S-CQ1---).

Group XX, claim(s) 1. (in part). 12(in part), ulawn to compounds of the formula (1) which constins a limiting of XI and XI and SX, claim(s). The O-PO(OH)-O or XI and XI into OPO(OH)-O or XI and XI into OPO(OH)-O or XI or X

Group XXI, claims 1(in part), 12(in part), drawn to compounds of formula (1) which consults a lithing of XI and X2 into OPO(OH)-O or X2 and X2 into OPO(OH)-NH- and at least one of XI or X2 or X2 in R1-Y1-A wherein R1 is not a harteropelic fing or or a sitrogen containing group and Y1 is (CH2), or O or C(O), A is a lith or (CH2), or O and Z1 is O (CH2), and O and Z1 is O (CH2), and O and Z1 is O (CH2), and O and Z2 is O or O(CH2)mi. Q1 and Q2 are "NR"-NR"R or at least one Q1 or Q2 is "NR"-NR'R'. (i.c. monophosphate exter-(HO),P(O)-O-CQ1--- or linked exter PO.Z2-P)

Groop XXII, claim(s) (tin part). 12(in part), drawn to compound of formula (t) which contains a linking of XI and X2 into d-PO(QH)-Off X2 and X3 into OPO(QH)-NH. and X3 and at least one of XI or X3 in XI in XI is the A party-cycle fing or a diverge containing upon part YI is (CH2), or O. or carborny'. A list or (CH2), or O and XI in NH. Z2 is O or O(CH2)m, Q1 and Q2 are "NR". NR'R" or at least one Q1 or Q2 is "NR". NR'R" (i.e. munophosphorus amilie ester -(HO),P(O)NH-CQ1--).

Group XXIII, claimit) (fin part). 12 (in part), chaven to compounds of the formula () which contains a liabing of XI and XZ into OPO(OBI-) or XZ and XZ into X

Group XXIV. claim(i) ((in part), Li2(in part), Grawn to compounds of the formula (())which coracins s lincking of XI and XI and XXIV. Car XI is RI-VI-A wherein RI is not a heterosystic ridg or or a threpton containing apon and Y is (CHZ), and O. or carboryl. A is s link or (CHZ), or O. and XI is (CH3), or CHARL ZI is 0 or OCCHZID, QI and Q2 are NR*-NR*R* or at least one Q1 or Q2 is NR*-NR*R* (i.e. phosphorms acid non an ener-(HO),P(O-CHZ-OCI)—).

Group XXV, claims 1(in part), 8(in part), 12(in part), drawn to compounds of formula (f) which consains a linking of XI and X2 into O-PO(OH)-O or X2 and X3 late O-PO(OH)-NH- and at least one of XI or X3 or X3 to R1-Y1-A wherein R1 is not a bacorogolic

fing or or a mirrogen containing group and Yi is NR2, A is a link or (CH2), or O and Zi is O (CH2), or O . Z2 is O of O(CH2)m, QI and Q2 are = NR* -NR*R* or at least one Q1 or Q2 is = NR* -NR*R*.

Porm PCT/ISA/210 (extra shoot) (July 1998)

INTERNATIONAL SEARCH REPORT

International application No. PCT/US01/08729

Group XXVI, claim(s) I(in pan), if in pan) If(in pan), drawn to compounds of formula (l) which contains a liching of XI and XI into O-PO(OH)-O or XI and XI into O-PO(OH)-NH- and all class one of XI or XI or XI or XI in R-IV-1-A wherein RI is an abscring-yield ring or or a nitrogen containing group and YI is NRZ, A is a link or (CHZ), or O and ZI is NR, ZZ is O or O(CHZ)m, QI and QZ an=NR'-NRZR' or at least one QI or QZ is =NR'-NRZR'. (i.e. monophesphorus amide ester -(HO) 3P(O)NH-CQ1---).

Group XXVII., claim(s) I (in part), 8 (in part) 12 (in part), drawn to compounds of the formula (I) which contains a limbing of X1 and X2 into O-PO(OH)-O or X2 and X3 into O-PO(OH)-NH- and X3 and Q2 are H2 or O. (i.e. monophosphorus amide ester -(HO),P(O)NH-CQ1---).

py XXVIII, claim(s) I(in part), 8 (in part), 12(in part), drawn to compounds of the formula (i) which contains a ed XI and XI into O-PO(OH)-O or XI and XI into O-PO(OH)-NH- and at

Whertin RI is not a heterocyclic ring or or a nitrogen comaining group and YI is (CH2) and O, or carbonyl . A is a link or (CH2), or O and ZI is S, Z2 is O or O(CH2)m, QI and Q2 are H2 or O, (i.e. monochiophosphaio seter -(HO),P(O)-S-CQ1---).

Group XXX, claim(s) I(in part). 12(in part), drawn to compounds of the formule (i) which contain X2and X3 linked as .NH-C(O)-NR1: and at least one of X1 or X3 or X3 is R1-Y1-A when in R1 is not a heterocyclic ring or or a nitrogen constaining group and Y1 is (CH3), or O, or carbonyl .A is a link or (CH3), or O and Z1 into O-PO(CH3). Or X2 and X3 into O-PO(CH3-NH-NH- and at least one of X1 or X2 or X3 is R1-Y1-A wherein R1 is not a heterocyclic ring or or a nitrogen containing group and Y1 is NR2. A is a link or (CH2), or O and Z1 is NR. Z2 is O or O(CH2)m. O1 and Q2 are H2

Group XXIV. exim(t) 1. (in part). 12(in part), drawn to compounds of the formula (l)which consints a laking of Xi and XX into O-PO(0H)-0 or XX and XX into O-PO(0H)-NH- and at least one of X (or of X2 or X3 is R1-Y1-A wherein R1 is not a heterowelle ring or or a through or constituing symp and Y1 is RR2, or extensy1. A is a link or (CH2), or O-200 XI is (CH2), or CR3H, Z2 is O or O/CH2)m. Q1 and Q2 srH2 or O. (i.e. phosphorus soid not an exter-

Group XXXI, claims 1(in part), 12(in part), drawn to compounds of Formula (1) which contain X2and X3 linked as - NH-C(O)-NR1- and at text one of X1 or X2 or X3 is R1-Y1-A wherein R1 is not a hateropylis ring or or a nirrogen containing group and Y1 is (CR12), or O or C(O). A is a link or (CR12), or O and Z1 is O (CR22), or O . Z2 is O or O(CR12), or O and Z1 is O (CR23), or O . Z2 is O or O(CR23), or O and Z1 is O (CR23), or O or C(D), P(O)-O-CQ1— or linked cuter PO-Z2-P)

Group XXXII, claim(s) 1(tin pan), 12(to pan), drawn to compounds of formula (t) which constin X2and X3 tidded as - NH-C(O)-NR1- and at least one of X1 or X2 or X3 is R1-Y1-A wherein R1 is not a heart-poyclic ring or or a alrogen containing group and Y1 is (CH2), or O, or arbony). A is a tide or (CH2), or O and Z1 is NH. Z2 is O or O(CH2)m. Q1 and Q2 are - NR*-NR** or at least one Q1 or Q2 is - NR*-NR**. (i.e. monophosphorus smitde ester -(HO),P(O)NH-CQ1--)

Group XXXIV. claim(s) 1(in part), 13(in part), drawn to campeneds of the formula (1) which contain X2aad X3 linked at -NH-C(O)-NR1- and at least one of X1 or X2 or X3 is R1-Y1-A. wherein R1 is not a bettroyed in fing or or a claimage containing group and Y1 is (CH3), or O creathoup, A is a link or (CH3), or O and Z1 is (CH3), or CR2H, Z2 is O or O(CH3)m Q1 and Q2 are -NR*-NR** or at least one Q1 or Q2 is -NR*-NR**. (i.e., phosphorus (HO),P(O)-S-CQ1---).

NHCQD-NRI- and at least one of XI or XI or XI is RI-YI-A whentis RI is not a betropytic ring or or a shirogen containing group and YI is NR2, AI = shire or (CH2), or O and ZI is O (CH3), or O . 22 is O or O(CH3)th QI and QI are =NR'-NR'R' or at least one QI or QI is =NR'-NR'R'. Oroup XXXV, claims 1(in part), 12(in part), drawn to compounds of formula (1) which comain X2and X3 linked as -

acid not an exter -(HO)2P(O-CH2-CQ1---).

Form PCT/ISA/210 (extra sheet) (July 1998) •

INTERNATIONAL SEARCH REPORT

International application No.

PCT/US01/08729

Group XXXVI, claims(s) It in part), 12(in part), drawn to compounds of formula (1) which consain X2 and X3 linked as -tNf-C(0)-NRI- and at least one of X1 or X2 or X3 is R1-Y1-A wherein R1 is not a heterosyclic ring or or a minogen containing group and Y1 is NR2, A is a link or (CH2), or 0-and Z1 is NH, Z2 is 0 or O(CH2)m, Q1 and Q2 are = NR*-tNF/N* or at least one Q1 or Q2 is = NR*-tNF/N*. (1.s. monophosphorus smide exter-tH0), P(0)NH-CQ1-

Group XXVVII, elizato) 1(in part), 12 (in part), drawn to compounds of the formula (!) which contain X2and X2 inlacts at .PH-C(O)-NR1- and X3 and at least one of X1 or X2 or X2 is R1-Y1-A whereth R1 is not a hactrocyclic ring or or a nitrogen containing proup and Y1 is MR2, A is a list or (CH2); or O and Z1 is S, Z is O or O(CH2)m, O1 and Q1 are = NR²-NR² or at least one Q1 or Q2 is = NR²-NR²! (i.e. manochiophosphate ester -(HO),P(O)-S-CQ1---).

Group XXXVIII. claim(s) ((in part), 12(in part), drawn to compounds of the formula (1) which consists a linking of X1 and X2 into 0-Po(OH)-O or X2 and X2 into 0-Po(OH)-NH- and at least one of X1 or X2 or X3 is R1-Y1-A wherein R1 is ton a heterocyclic ring or a nitrogen containing group and Y1 is NR2, or carboxy', A is a link or (CH2), or O and Z1 is (CH3), or CR2H, Z2 is O or O(CH2)m, Q1 and Q2 or = NR*-NR*N* or at least one Q1 or Q2 is = NR*-NR*N*.

Groups XXXIX-LXXVI are the same as the above 1-XXXXII energy that RI is a heterocycle or a or a mirogen donor. These will be the same civities (in part) as those above.

Group LXXVII claim I (in part) and I2(in part) any compounds not found in Groups I-LXXVII

Group LXXVIII, ciaims 13-22. Urawn to a method of inhibiting LPA activity on an LPA receptor using compounds of

Group LXXIX, claims 23-26, drawn to a method of treating cancer.

Group LXXX, claims 27-30, drawn to a method of cohancing cell proliferation

Group LXXXI, claims 31-33, drawn to a method of treating a would

Groups LXXXII claim 34, process for the preparation of phosphorus containing compounds

The inventions listed as Groups I-LXXXII do not relate to a single inventive concept under PCT Rule 13.1 because,

constitute a large structural common core since there are hundreds of thousands of groups which contain a -CH-, it is also noted that the -CH- is not a small structural portion which is applicant's contribution to the art. The claims contain estern, acid, amidea, thicestern, heterocyclic phosphorus and non phosphorus containing groups which only have a -CH- as the common core. Therefore there is no special technical feature unting all the various compounds. under PCT Rule 13.2, they lack the same or corresponding special technical features for the following reasons: Group I has a series of various senseures which only contains the common core -CH-. The -CH- does not

compounds and the various methods of using the compounds since there are known compounds being claimed such as those compounds found in CA:116:230686, CA:117:166460,CA:83:10823, CA:117:1602002 or 115:200251. These compounds have their own various united and their various methods of being prepared. Therefore there is no unity of invention since there is no special technical feature which unites the groups by providing a compound; a method specially designed to prepare the compound and a method of using the compound that is specially linked to by a special Also there is no special technical feature among the cumpounds being claimed and the method of preparing the

Form PCT/ISA/210 (extra sheet) (July 1998) *